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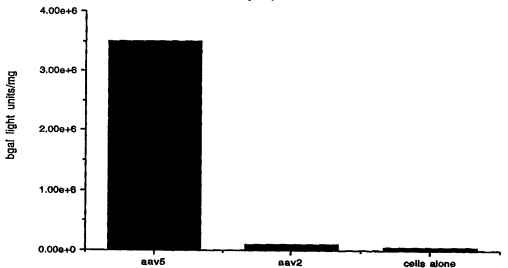
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(54) Title: AAV5 VECTOR AND USES THEREOF

#### Apical transduction of human airway epithelia with rAAV2 and rAAV5



#### (57) Abstract

The present invention provides an adeno-associated virus 5 (AAV5) virus and vectors and particles derived therefrom. In addition, the present invention provides methods of delivering a nucleic acid to a cell using the AAV5 vectors and particles.

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### **AAV5 VECTOR AND USES THEREOF**

This application claims priority to U.S. provisional application Serial No. 60/087029 filed on May 28, 1998. The 60/087029 provisional patent application is herein incorporated by this reference in its entirety.

### **BACKGROUND OF THE INVENTION**

#### Field of the Invention

The present invention provides adeno-associated virus 5 (AAV5) and vectors derived therefrom. Thus, the present invention relates to AAV5 vectors for and methods of delivering nucleic acids to cells of subjects.

### **Background Art**

Adeno associated virus (AAV) is a small nonpathogenic virus of the parvoviridae family (for review see 28). AAV is distinct from the other members of this family by its dependence upon a helper virus for replication. In the absence of a helper virus, AAV has been shown to integrate in a locus specific manner into the q arm of chromosome 19 (21). The approximately 5 kb genome of AAV consists of one segment of single stranded DNA of either plus or minus polarity. Physically, the parvovirus virion is non-enveloped and its icosohedral capsid is approximately 20-25 nm in diameter.

isolated from humans or primates and are referred to as AAV types 1-6 (1). The most extensively studied of these isolates is AAV type 2 (AAV2). The genome of AAV2 is 4680 nucleotides in length and contains two open reading frames (ORFs), the right ORF and the left ORF. The left ORF encodes the non-structural Rep proteins, Rep40, Rep52, Rep68 and Rep78, which are involved in regulation of replication and transcription in addition to the production of single-stranded progeny genomes (5-8, 11, 12, 15, 17, 19, 21-23, 25, 34, 37-40). Furthermore, two of the Rep proteins have been associated with the preferential integration of AAV genomes into a region of the q arm

of human chromosome 19. Rep68/78 have also been shown to possess NTP binding activity as well as DNA and RNA helicase activities. The Rep proteins possess a nuclear localization signal as well as several potential phosphorylation sites. Mutation of one of these kinase sites resulted in a loss of replication activity.

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The ends of the genome are short inverted terminal repeats which have the potential to fold into T-shaped hairpin structures that serve as the origin of viral DNA replication. Within the ITR region two elements have been described which are central to the function of the ITR, a GAGC repeat motif and the terminal resolution site (TRS). The repeat motif has been shown to bind Rep when the ITR is in either a linear or hairpin conformation (7, 8, 26).

This binding serves to position Rep68/78 for cleavage at the TRS which occurs in a site- and strand-specific manner. In addition to their role in replication, these two elements appear to be central to viral integration. Contained within the chromosome 19 integration locus is a Rep binding site with an adjacent TRS. These elements have been shown to be functional and necessary for locus specific integration.

The AAV2 virion is a non-enveloped, icosohedral particle approximately 20-25 nm in diameter. The capsid is composed of three related proteins referred to as VP1,2 and 3 which are encoded by the right ORF. These proteins are found in a ratio of 1:1:10 respectively. The capsid proteins differ from each other by the use of alternative splicing and an unusual start codon. Deletion analysis of has shown that removal or alteration of AAV2 VP1 which is translated from an alternatively spliced message results in a reduced yield of infections particles (15, 16, 38). Mutations within the VP3 coding region result in the failure to produce any single-stranded progeny DNA or infectious particles (15, 16, 38).

The following features of the characterized AAVs have made them attractive vectors for gene transfer (16). AAV vectors have been shown *in vitro* to stably integrate into the cellular genome; possess a broad host range; transduce both dividing and non dividing cells *in vitro* and *in vivo* (13, 20, 30, 32) and maintain high levels of

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expression of the transduced genes (41). Viral particles are heat stable, resistant to solvents, detergents, changes in pH, temperature, and can be concentrated on CsCl gradients (1,2). Integration of AAV provirus is not associated with any long term negative effects on cell growth or differentiation (3,42). The ITRs have been shown to be the only cis elements required for replication, packaging and integration (35) and may contain some promoter activities (14).

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AAV2 was originally thought to infect primate and non-primate cell types provided the appropriate helper virus was present. However, the inability of AAV2 to infect certain cell types is now known to be due to the particular cellular tropism exhibited by the AAV2 virus. Recent work has shown that some cell lines are transduced very poorly by AAV2 (30). Binding studies have indicated that heparin sulfate proteoglycans are necessary for high efficiency transduction with AAV2. AAV5 is a unique member of the parvovirus family. The present DNA hybridization data indicate a low level of homology with the published AAV1-4 sequences (31). The present invention shows that, unlike AAV2, AAV5 transduction is not effected by heparin as AAV2 is and therefore will not be restricted to the same cell types as AAV2.

The present invention provides a vector comprising the AAV5 virus or a vector comprising subparts of the virus, as well as AAV5 viral particles. While AAV5 is similar to AAV2, the two viruses are found herein to be physically and genetically distinct. These differences endow AAV5 with some unique properties and advantages which better suit it as a vector for gene therapy. For example, one of the limiting features of using AAV2 as a vector for gene therapy is production of large amounts of virus. Using standard production techniques, AAV5 is produced at a 10-50 fold higher level compared to AAV2. Because of its unique TRS site and rep proteins, AAV5 should also have a distinct integration locus compared to AAV2.

Furthermore, as shown herein, AAV5 capsid protein, again surprisingly, is

distinct from AAV2 capsid protein and exhibits different tissue tropism, thus making
AAV5 capsid-containing particles suitable for transducing cell types for which AAV2
is unsuited or less well-suited. AAV2 and AAV5 have been shown to be serologically

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distinct and thus, in a gene therapy application, AAV5, and AAV5-derived vectors, would allow for transduction of a patient who already possess neutralizing antibodies to AAV2 either as a result of natural immunological defense or from prior exposure to AAV2 vectors. Another advantage of AAV5 is that AAV5 cannot be rescued by other serotypes. Only AAV5 can rescue the integrated AAV5 genome and effect replication, thus avoiding unintended replication of AAV5 caused by other AAV serotypes. Thus, the present invention, by providing these new recombinant vectors and particles based on AAV5 provides a new and highly useful series of vectors.

#### 10 SUMMARY OF THE INVENTION

The present invention provides a nucleic acid vector comprising a pair of adenoassociated virus 5 (AAV5) inverted terminal repeats and a promoter between the inverted terminal repeats.

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The present invention further provides an AAV5 particle containing a vector comprising a pair of AAV2 inverted terminal repeats.

Additionally, the instant invention provides an isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO:1 (AAV5 genome). Furthermore, the present invention provides an isolated nucleic acid consisting essentially of the nucleotide sequence set forth in SEQ ID NO:1 (AAV5 genome).

25 protein, for example, the nucleic acid as set forth in SEQ ID NO:10. Additionally provided is an isolated full-length AAV5 Rep protein or a unique fragment thereof. Additionally provided is an isolated AAV5 Rep 40 protein having the amino acid sequence set forth in SEQ ID NO:12, or a unique fragment thereof. Additionally provided is an isolated AAV5 Rep 52 protein having the amino acid sequence set forth in SEQ ID NO:2, or a unique fragment thereof. Additionally provided is an isolated AAV5 Rep 52 protein having the amino acid sequence set forth AAV5 Rep 68 protein, having the amino acid sequence set forth in SEQ ID NO:14 or a unique fragment thereof. Additionally provided is an isolated AAV5 Rep 78 protein

having the amino acid sequence set forth in SEQ ID NO:3, or a unique fragment thereof. The sequences for these proteins are provided below in the Sequence Listing and elsewhere in the application where the proteins are described.

The present invention further provides an isolated AAV5 capsid protein, VP1, having the amino acid sequence set forth in SEQ ID NO:4, or a unique fragment thereof. Additionally provided is an isolated AAV5 capsid protein, VP2, having the amino acid sequence set forth in SEQ ID NO:5, or a unique fragment thereof. Also provided is an isolated AAV5 capsid protein, VP3, having the amino acid sequence set forth in SEQ ID NO:6, or a unique fragment thereof.

The present invention additionally provides an isolated nucleic acid encoding AAV5 capsid protein, for example, the nucleic acid set forth in SEQ ID NO:7, or a unique fragment thereof.

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The present invention further provides an AAV5 particle comprising a capsid protein consisting essentially of the amino acid sequence set forth in SEQ ID NO:4, or a unique fragment thereof.

Additionally provided by the present invention is an isolated nucleic acid comprising an AAV5 p5 promoter having the nucleic acid sequence set forth in SEQ ID NO:18, or a unique fragment thereof.

The instant invention provides a method of screening a cell for infectivity by

AAV5 comprising contacting the cell with AAV5 and detecting the presence of AAV5 in the cells.

The present invention further provides a method of delivering a nucleic acid to a cell comprising administering to the cell an AAV5 particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to the cell.

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The present invention also provides a method of delivering a nucleic acid to a subject comprising administering to a cell from the subject an AAV5 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, and returning the cell to the subject, thereby delivering the nucleic acid to the subject.

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The present invention also provides a method of delivering a nucleic acid to a cell in a subject comprising administering to the subject an AAV5 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to a cell in the subject.

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The instant invention further provides a method of delivering a nucleic acid to a cell in a subject having antibodies to AAV2 comprising administering to the subject an AAV5 particle comprising the nucleic acid, thereby delivering the nucleic acid to a cell in the subject.

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### BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 shows Heparin inhibition results. Cos cells were plated in 12 well dishes at 5X10<sup>4</sup> cells per well. Serial dilutions of AAV2 or AAV5 produced and purified as previously described and supplemented with 5X10<sup>5</sup> particles of wt adenovirus were incubated for 1 hr at Rt in the presence of 20 μg/ml heparin (sigma). Following this incubation the virus was added to the cells in 400 μl of media for 1 hr after which the media was removed, the cells rinsed and fresh media added. After 24 hrs the plates were stained for Bgal activity.

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Figure 2 shows AAV2 and AAV5 vector and helper complementation. Recombinant AAV particles were produced as previously described using a variety of vector and helper plasmids as indicated the bottom of the graph. The vector plasmids contained the Bgal gene with and RSV promoter and flanked by either AAV2 ITRs (2ITR) or AAV5 ITRs (5ITR). The helper plasmids tested contained either AAV2 Rep and cap genes (2repcap) AAV5 rep and cap genes with or without an SV40 promoter (5repcapA and 5repcapb respectively) only the AAV2 rep gene (2rep) in varying

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amounts (1) or (.5) or an empty vector (pUC). The resulting AAV particles were then titered on cos cells. AAV particles were only produced when the same serotype of ITR and Rep were present.

Figure 3 shows AAV2 and AAV5 tissue tropism. Transduction of a variety of cell types indicated that AAV2 and AAV5 transduce cells with different efficiencies. Equal number of either AAV2 or AAV5 particles were used to transduce a variety of cell types and the number of bgal positive cells is reported.

Figure 4 is a sequence comparison of the AAV2 genome and the AAV5 genome.

Figure 5 is a sequence comparison of the AAV2 VP1 capsid protein and the AAV5 VP1 capsid protein.

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Figure 6 is a sequence comparison of the AAV2 rep 78 protein and the AAV5 rep 78 protein.

Figure 7 shows the transduction of airway epithelial cells by AAV5. Primary airway epithelial cells were cultured and plated. Cells were transducted with an equivalent number of rAAV2 or rAAV5 particles containing a nuclear localized β-gal transgene with 50 particles of virus/cell (MOI 50) and continued in culture for 10 days. β-gal activity was determined and the relative transduction efficiency compared. AAV5 transduced these cells 50- fold more efficiently than AAV2. This is the first time apical cells or cells exposed to the air have been shown to be infected by a gene therapy agent.

Figure 8 shows transduction of striated muscle by AAV5. Chicken myoblasts were cultured and plated. Cells were allowed to fuse and then transduced with a similar number of particles of rAAV2 or rAAV5 containing a nuclear localized  $\beta$ -gal transgene after 5 days in culture. The cells were stained for  $\beta$ -gal activity and the relative

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transduction efficiency compared. AAV5 transduced these cells approximately 16 fold more efficiently than AAV2.

Figure 9 shows transduction of rat brain explants by AAV5. Primary neonatal rat brain explants were prepared. After 7 days in culture, cells were transduced with a similar number of particles of rAAV5 containing a nuclear localized β-gal transgene. After 5 days in culture, the cells were stained for β-gal activity. Transduction was detected in a variety of cell types including astrocytes, neuronal cells and glial cells.

Figure 10 shows transduction of human umbilical vein endothelial cells by AAV5. Human umbilical vein endothelial cells were cultured and plated. Cells were transduced with rAAV2 or rAAV5 containing a nuclear localized β-gal transgene with 10 particles of virus/ cell (MOI 5) in minimal media then returned to complete media. After 24 hrs in culture, the cells were stained for β-gal activity and the relative transduction efficiency compared. As shown in AAV5 transduced these cell 5-10 fold more efficiently than AAV2.

### DETAILED DESCRIPTION OF THE INVENTION

As used in the specification and in the claims, "a" can mean one or more, depending upon the context in which it is used. The terms "having" and "comprising" are used interchangeably herein, and signify open ended meaning.

The present application provides a recombinant adeno-associated virus 5

(AAV5). This virus has one or more of the characteristics described below. The compositions of the present invention do not include wild-type AAV5. The methods of the present invention can use either wild-type AAV5 or recombinant AAV5-based delivery.

The present invention provides novel AAV5 particles, recombinant AAV5 vectors, recombinant AAV5 virions and novel AAV5 nucleic acids and polypeptides.

An AAV5 particle is a viral particle comprising an AAV5 capsid protein. A

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recombinant AAV5 vector is a nucleic acid construct that comprises at least one unique nucleic acid of AAV5. A recombinant AAV5 virion is a particle containing a recombinant AAV5 vector, wherin the particle can be either an AAV5 particle as described herein or a non-AAV5 particle. Alternatively, the recombinant AAV5 virion is an AAV5 particle containing a recombinant vector, wherein the vector can be either an AAV5 vector as described herein or a non-AAV5 vector. These vectors, particles, virions, nucleic acids and polypeptides are described below.

The present invention provides the nucleotide sequence of the AAV5 genome and vectors and particles derived therefrom. Specifically, the present invention 10 provides a nucleic acid vector comprising a pair of AAV5 inverted terminal repeats (ITRs) and a promoter between the inverted terminal repeats. While the rep proteins of AAV2 and AAV5 will bind to either a type 2 ITR or a type 5 ITR, efficient genome replication only occurs when type 2 Rep replicates a type 2 ITR and a type 5 Rep replicates a type 5 ITR. This specificity is the result of a difference in DNA cleavage 15 specificity of the two Reps which is necessary for replication. AAV5 Rep cleaves at CGGT^GTGA (SEQ ID NO: 21) and AAV2 Rep cleaves at CGGT^TGAG (SEQ ID NO: 22) (Chiorini et al., 1999. J. Virol. 73 (5) 4293-4298). Mapping of the AAV5 ITR terminal resolution site (TRS) identified this distinct cleavage site, CGGT^GTGA, 20 which is absent from the ITRs of other AAV serotypes. Therefore, the minimum sequence necessary to distinguish AAV5 from AAV2 is the TRS site where Rep cleaves in order to replicate the virus. Examples of the type 5 ITRs are shown in SEQ ID NO: 19 and SEQ ID NO: 20, AAV5 ITR "flip" and AAV5 "flop", respectively. Minor modifications in an ITR of either orientation are contemplated and are those that 25 will not interfere with the hairpin structure formed by the AAV5 ITR as described herein and known in the art. Furthermore, to be considered within the term "AAV5 ITR" the nucleotide sequence must retain one or more features described herein that distinguish the AAV5 ITR from the ITRs of other serotypes, e.g. it must retain the Rep binding site described herein.

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The D- region of the AAV5 ITR (SEQ ID NO: 23), a single stranded region of the ITR, inboard of the TRS site, has been shown to bind a factor which depending on

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its phosphorylation state correlates with the conversion of the AAV from a single stranded genome to a transcriptionally active form that allows for expression of the viral DNA. This region is conserved between AAV2, 3, 4,and 6 but is divergent in AAV5. The D+ region is the reverse complement of the D- region.

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The promoter can be any desired promoter, selected by known considerations, such as the level of expression of a nucleic acid functionally linked to the promoter and the cell type in which the vector is to be used. That is, the promoter can be tissue/cellspecific. Promoters can be prokaryotic, eukaryotic, fungal, nuclear, mitochondrial, viral or plant promoters. Promoters can be exogenous or endogenous to the cell type being transduced by the vector. Promoters can include, for example, bacterial promoters, known strong promoters such as SV40 or the inducible metallothionein promoter, or an AAV promoter, such as an AAV p5 promoter. Additionally, chimeric regulatory promoters for targeted gene expression can be utilized. Examples of these regulatory systems, which are known in the art, include the tetracycline based regulatory system which utilizes the tet transactivator protein (tTA), a chimeric protein containing the VP16 activation domain fused to the tet repressor of Escherichia coli, the IPTG based regulatory system, the CID based regulatory system, and the Ecdysone based regulatory system (44). Other promoters include promoters derived from actin genes, immunoglobulin genes, cytomegalovirus (CMV), adenovirus, bovine papilloma virus, adenoviral promoters, such as the adenoviral major late promoter, an inducible heat shock promoter, respiratory syncytial virus, Rous sarcomas virus (RSV), etc., specifically, the promoter can be AAV2 p5 promoter or AAV5 p5 promoter. More specifically, the AAV5 p5 promoter can be about same location in SEQ ID NO: 1 as the AAV2 p5 promoter, in the corresponding AAV2 published sequence. Additionally, the p5 promoter may be enhanced by nucleotides 1-130 of SEQ ID NO:1. Furthermore, smaller fragments of p5 promoter that retain promoter activity can readily be determined by standard procedures including, for example, constructing a series of deletions in the p5 promoter, linking the deletion to a reporter gene, and determining whether the reporter gene is expressed, i.e., transcribed and/or translated. The promoter can be the promoter of any of the AAV serotypes, and can be the p19 promoter (SEQ ID NO: 16) or the p40 promoter set forth in the sequence listing as SEQ ID NO: 17.

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It should be recognized that any errors in any of the nucleotide sequences disclosed herein can be corrected, for example, by using the hybridization procedure described below with various probes derived from the described sequences such that the coding sequence can be reisolated and resequenced. Rapid screening for point mutations can also be achieved with the use of polymerase chain reaction-single strand conformation polymorphism (PCR-SSCP) (43). The corresponding amino acid sequence can then be corrected accordingly.

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10 nucleic acid functionally linked to the promoter. By "heterologous nucleic acid" is meant that any heterologous or exogenous nucleic acid, i.e. not normally found in wild-type AAV5 can be inserted into the vector for transfer into a cell, tissue or organism. By "functionally linked" is meant that the promoter can promote expression of the heterologous nucleic acid, as is known in the art, and can include the appropriate orientation of the promoter relative to the heterologous nucleic acid. Furthermore, the heterologous nucleic acid preferably has all appropriate sequences for expression of the nucleic acid. The nucleic acid can include, for example, expression control sequences, such as an enhancer, and necessary information processing sites, such as ribosome binding sites, RNA splice sites, polyadenylation sites, and transcriptional terminator sequences.

The heterologous nucleic acid can encode beneficial proteins or polypeptides that replace missing or defective proteins required by the cell or subject into which the vector is transferred or can encode a cytotoxic polypeptide that can be directed, *e.g.*, to cancer cells or other cells whose death would be beneficial to the subject. The heterologous nucleic acid can also encode antisense RNAs that can bind to, and thereby inactivate, mRNAs made by the subject that encode harmful proteins. The heterologous nucleic acid can also encode ribozymes that can effect the sequence-specific inhibition of gene expression by the cleavage of mRNAs. In one embodiment, antisense polynucleotides can be produced from a heterologous expression cassette in an AAV5 vector construct where the expression cassette contains a sequence that promotes cell-type specific expression (Wirak *et al.*, *EMBO* 10:289

(1991)). For general methods relating to antisense polynucleotides, see *Antisense RNA* and *DNA*, D. A. Melton, Ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, NY (1988).

5 Examples of heterologous nucleic acids which can be administered to a cell or subject as part of the present AAV5 vector can include, but are not limited to the following: nucleic acids encoding secretory and nonsecretory proteins, nucleic acids encoding therapeutic agents, such as tumor necrosis factors (TNF), such as TNF-α; interferons, such as interferon- $\alpha$ , interferon- $\beta$ , and interferon- $\gamma$ ; interleukins, such as IL-1, IL-1β, and ILs -2 through -14; GM-CSF; adenosine deaminase; cellular growth 10 factors, such as lymphokines; soluble CD4; Factor VIII; Factor IX; T-cell receptors; LDL receptor; ApoE; ApoC; alpha-1 antitrypsin; ornithine transcarbamylase (OTC); cystic fibrosis transmembrane receptor (CFTR); insulin; Fc receptors for antigen binding domains of antibodies, such as immunoglobulins; anit-HIV decoy tar elements; 15 and antisense sequences which inhibit viral replication, such as antisense sequences which inhibit replication of hepatitis B or hepatitis non-A, non-B virus. The nucleic acid is chosen considering several factors, including the cell to be transfected. Where the target cell is a blood cell, for example, particularly useful nucleic acids to use are those which allow the blood cells to exert a therapeutic effect, such as a gene encoding 20 a clotting factor for use in treatment of hemophilia. Another target cell is the lung airway cell, which can be used to administer nucleic acids, such as those coding for the cystic fibrosis transmembrane receptor, which could provide a gene therapeutic treatment for cystic fibrosis. Other target cells include muscle cells where useful nucleic acids, such as those encoding cytokines and growth factors, can be transduced and the protein the nucleic acid encodes can be expressed and secreted to exert its 25 effects on other cells, tissues and organs, such as the liver. Furthermore, the nucleic acid can encode more than one gene product, limited only, if the nucleic acid is to be packaged in a capsid, by the size of nucleic acid that can be packaged.

Furthermore, suitable nucleic acids can include those that, when transferred into a primary cell, such as a blood cell, cause the transferred cell to target a site in the body where that cell's presence would be beneficial. For example, blood cells such as TIL

cells can be modified, such as by transfer into the cell of a Fab portion of a monoclonal antibody, to recognize a selected antigen. Another example would be to introduce a nucleic acid that would target a therapeutic blood cell to tumor cells. Nucleic acids useful in treating cancer cells include those encoding chemotactic factors which cause an inflammatory response at a specific site, thereby having a therapeutic effect.

Cells, particularly blood cells, muscle cells, airway epithelial cells, brain cells and endothelial cells having such nucleic acids transferred into them can be useful in a variety of diseases, syndromes and conditions. For example, suitable nucleic acids include nucleic acids encoding soluble CD4, used in the treatment of AIDS and  $\alpha$ -antitrypsin, used in the treatment of emphysema caused by  $\alpha$ -antitrypsin deficiency. Other diseases, syndromes and conditions in which such cells can be useful include, for example, adenosine deaminase deficiency, sickle cell deficiency, brain disorders such as Alzheimer's disease, thalassemia, hemophilia, diabetes, phenylketonuria, growth disorders and heart diseases, such as those caused by alterations in cholesterol metabolism, and defects of the immune system.

As another example, hepatocytes can be transfected with the present vectors having useful nucleic acids to treat liver disease. For example, a nucleic acid encoding OTC can be used to transfect hepatocytes ( $ex\ vivo$  and returned to the liver or  $in\ vivo$ ) to treat congenital hyperammonemia, caused by an inherited deficiency in OTC. Another example is to use a nucleic acid encoding LDL to target hepatocytes  $ex\ vivo$  or  $in\ vivo$  to treat inherited LDL receptor deficiency. Such transfected hepatocytes can also be used to treat acquired infectious diseases, such as diseases resulting from a viral infection. For example, transduced hepatocyte precursors can be used to treat viral hepatitis, such as hepatitis B and non-A, non-B hepatitis, for example by transducing the hepatocyte precursor with a nucleic acid encoding an antisense RNA that inhibits viral replication. Another example includes transferring a vector of the present invention having a nucleic acid encoding a protein, such as  $\alpha$ -interferon, which can confer resistance to the hepatitis virus.

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For a procedure using transfected hepatocytes or hepatocyte precursors, hepatocyte precursors having a vector of the present invention transferred in can be grown in tissue culture, removed from the tissue culture vessel, and introduced to the body, such as by a surgical method. In this example, the tissue would be placed directly into the liver, or into the body cavity in proximity to the liver, as in a transplant or graft. Alternatively, the cells can simply be directly injected into the liver, into the portal circulatory system, or into the spleen, from which the cells can be transported to the liver via the circulatory system. Furthermore, the cells can be attached to a support, such as microcarrier beads, which can then be introduced, such as by injection, into the peritoneal cavity. Once the cells are in the liver, by whatever means, the cells can then express the nucleic acid and/or differentiate into mature hepatocytes which can express the nucleic acid.

The AAV5-derived vector can include any normally occurring AAV5 sequences in addition to an ITR and promoter. Examples of vector constructs are provided below.

The present vector or AAV5 particle or recombinant AAV5 virion can utilize any unique fragment of these present AAV5 nucleic acids, including the AAV5 nucleic acids set forth in SEQ ID NOS: 1 and 7-11, 13, 15, 16, 17, and 18. To be unique, the fragment must be of sufficient size to distinguish it from other known sequences, most readily determined by comparing any nucleic acid fragment to the nucleotide sequences of nucleic acids in computer databases, such as GenBank. Such comparative searches are standard in the art. Typically, a unique fragment useful as a primer or probe will be at least about 8 or 10, preferable at least 20 or 25 nucleotides in length, depending upon the specific nucleotide content of the sequence. Additionally, fragments can be, for example, at least about 30, 40, 50, 75, 100, 200 or 500 nucleotides in length and can encode polypeptides or be probes. The nucleic acid can be single or double stranded, depending upon the purpose for which it is intended. Where desired, the nucleic acid can be RNA.

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The present invention further provides an AAV5 capsid protein to contain the vector. In particular, the present invention provides not only a polypeptide comprising

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all three AAV5 coat proteins, i.e., VP1, VP2 and VP3, but also a polypeptide comprising each AAV5 coat protein individually, SEQ ID NOS: 4, 5, and 6. respectively. Thus an AAV5 particle comprising an AAV5 capsid protein comprises at least one AAV5 coat protein VP1, VP2 or VP3. An AAV5 particle comprising an AAV5 capsid protein can be utilized to deliver a nucleic acid vector to a cell, tissue or 5 subject. For example, the herein described AAV5 vectors can be encapsidated in an AAV5 capsid-derived particle and utilized in a gene delivery method. Furthermore. other viral nucleic acids can be encapsidated in the AAV5 particle and utilized in such delivery methods. For example, an AAV1, 2,3,4,or 6 vector (e.g. AAV1,2,3,4,or 6 ITR and nucleic acid of interest )can be encapsidated in an AAV5 particle and 10 administered. Furthermore, an AAV5 chimeric capsid incorporating both AAV2 capsid and AAV5 capsid sequences can be generated, by standard cloning methods, selecting regions from the known sequences of each protein as desired. For example, particularly antigenic regions of the AAV2 capsid protein can be replaced with the corresponding region of the AAV5 capsid protein. In addition to chimeric capsids incorporating 15 AAV2 capsid sequences, chimeric capsids incorporating AAV1, 3, 4, or 6 and AAV5 capsid sequences can be generated, by standard cloning methods, selecting regions from the known sequences of each protein as desired.

The capsids can also be modified to alter their specific tropism by genetically altering the capsid to encode a specific ligand to a cell surface receptor. Alternatively, the capsid can be chemically modified by conjugating a ligand to a cell surface receptor. By genetically or chemically altering the capsids, the tropism can be modified to direct AAV5 to a particular cell or population of cells. The capsids can also be altered immunologically by conjugating the capsid to an antibody that recognizes a specific protein on the target cell or population of cells.

The capsids can also be assembled into empty particles by expression in mammalian, bacterial, fungal or insect cells. For example, AAV2 particles are known to be made from VP3 and VP2 capsid proteins in baculovirus. The same basic protocol can produce an empty AAV5 particle comprising an AAV5 capsid protein.

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The herein described recombinant AAV5 nucleic acid derived vector can be encapsidated in an AAV particle. In particular, it can be encapsidated in an AAV1 particle, an AAV2 particle, an AAV3 particle, an AAV4 particle, an AAV5 particle or an AAV6 particle, a portion of any of these capsids, or a chimeric capsid particle as described above, by standard methods using the appropriate capsid proteins in the encapsidation process, as long as the nucleic acid vector fits within the size limitation of the particle utilized. The encapsidation process itself is standard in the art. The AAV5 replication machinery, i.e. the rep initiator proteins and other functions required for replication, can be utilized to produce the AAV5 genome that can be packaged in an AAV1, 2, 3, 4, 5 or 6 capsid.

The recombinant AAV5 virion containing a vector can also be produced by recombinant methods utilizing multiple plasmids. In one example, the AAV5 rep nucleic acid would be cloned into one plasmid, the AAV5 ITR nucleic acid would be cloned into another plasmid and the AAV1, 2, 3, 4, 5 or 6 capsid nucleic acid would be cloned on another plasmid. These plasmids would then be introduced into cells. The cells that were efficiently transduced by all three plasmids, would exhibit specific integration as well as the ability to produce AAV5 recombinant virus. Additionally, two plasmids could be used where the AAV5 rep nucleic acid would be cloned into one plasmid and the AAV5 ITR and AAV5 capsid would be cloned into another plasmid. These plasmids would then be introduced into cells. The cells that were efficiently transduced by both plasmids, would exhibit specific integration as well as the ability to produce AAV5 recombinant virus.

An AAV5 capsid polypeptide encoding the entire VP1, VP2, and VP3 polypeptide can overall has greater than 56% homology to the polypeptide having the amino acid sequence encoded by nucleotides in SEQ ID NOS:7,8 and 9, as shown in figures 4 and 5. The capsid protein can have about 70% homology, about 75% homology, 80% homology, 85% homology, 90% homology, 95% homology, 98% homology, 99% homology, or even 100% homology to the protein having the amino acid sequence encoded by the nucleotides set forth in SEQ ID NOS:7, 8 or 9. The percent homology used to identify proteins herein, can be based on a nucleotide-by-

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nucleotide comparison or more preferable is based on a computerized algorithm as described herein. Variations in the amino acid sequence of the AAV5 capsid protein are contemplated herein, as long as the resulting particle comprising an AAV5 capsid protein remains antigenically or immunologically distinct from AAV1, AAV2, AAV3,

5 AAV4 or AAV6 capsid, as can be routinely determined by standard methods.

Specifically, for example, ELISA and Western blots can be used to determine whether a viral particle is antigenically or immunologically distinct from AAV2 or the other serotypes. Furthermore, the AAV5 particle preferably retains tissue tropism distinction from AAV2, such as that exemplified in the examples herein. An AAV5 chimeric particle comprising at least one AAV5 coat protein may have a different tissue tropism from that of an AAV5 particle consisting only of AAV5 coat proteins, but is still distinct from the tropism of an AAV2 particle.

The invention further provides a recombinant AAV5 virion, comprising an AAV5 particle containing, *i.e.*, encapsidating, a vector comprising a pair of AAV5 inverted terminal repeats. The recombinant vector can further comprise an AAV5 Repencoding nucleic acid. The vector encapsidated in the particle can further comprise an exogenous nucleic acid inserted between the inverted terminal repeats. AAV5 Rep confers targeted integration and efficient replication, thus production of recombinant AAV5, comprising AAV5 Rep, yields more particles than production of recombinant AAV2. Since AAV5 is more efficient at replicating and packaging its genome, the exogenous nucleic acid inserted, or in the AAV5 capsids of the present invention, between the inverted terminal repeats can be packaged in the AAV1, 2, 3, 4, or 6 capsids to achieve the specific tissue tropism conferred by the capsid proteins.

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The invention further contemplates chimeric recombinant ITRs that contains a rep binding site and a TRS site recognized by that Rep protein. By "Rep protein" is meant all four of the Rep proteins, Rep 40, Rep 78, Rep 52, Rep 68. Alternatively, "Rep protein" could be one or more of the Rep proteins described herein. One example of a chimeric ITR would consist of an AAV5 D region (SEQ ID NO: 23), an AAV5 TRS site (SEQ ID NO: 21), an AAV2 hairpin and an AAV2 binding site. Another example would be an AAV5 D region, an AAV5 TRS site, an AAV3 hairpin and an

AAV3 binding site. In these chimeric ITRs, the D region can be from AAV1, 2, 3, 4, 5 or 6. The hairpin can be derived from AAV 1, 2, 3, 4, 5, 6. The binding site can be derived from any of AAV1, 2, 3, 4, 5 or 6. Preferably, the D region and the TRS are from the same serotype.

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The chimeric ITRs can be combined with AAV5 Rep protein and any of the AAV serotype capsids to obtain recombinant virion. For example, recombinant virion can be produced by an AAV5 D region, an AAV5 TRS site, an AAV2 hairpin, an AAV2 binding site, AAV5 Rep protein and AAV1 capsid. This recombinant virion would possess the cellular tropism conferred by the AAV1 capsid protein and would possess the efficient replication conferred by the AAV5 Rep.

Other examples of the ITR, Rep protein and Capsids that will produce recombinant virus are provided in the list below:

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5ITR + 5Rep + 5Cap=virus
5ITR + 5Rep + 1Cap=virus
5ITR + 5Rep + 2Cap=virus
5ITR + 5Rep + 2Cap=virus
5ITR + 5Rep + 3Cap=virus
20 5ITR + 5Rep + 4Cap=virus
5ITR + 5Rep + 6Cap=virus
1ITR + 1Rep + 5Cap=virus
2ITR + 2Rep + 5Cap=virus
3ITR + 3Rep + 5Cap=virus
4ITR + 4Rep + 5Cap=virus
6ITR + 6Rep + 5Cap=virus
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In any of the constructs described herein, inclusion of a promoter is preferred. As used in the constructs herein, unless otherwise specified, Cap (capsid) refers to any of AAV5 VP1, AAV5 VP2, AAV5 VP3, combinations thereof, functional fragments of any of VP1, VP2 or VP3, or chimeric capsids as described herein. The ITRs of the

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constructs described herein, can be chimeric recombinant ITRs as described elsewhere in the application.

Conjugates of recombinant or wild-type AAV5 virions and nucleic acids or proteins can be used to deliver those molecules to a cell. For example, the purified AAV5 can be used as a vehicle for delivering DNA bound to the exterior of the virus. Examples of this are to conjugate the DNA to the virion by a bridge using poly-L-lysine or other charged molecule. Also contemplated are virosomes that contain AAV5 structural proteins (AAV5 capsid proteins), lipids such as DOTAP, and nucleic acids that are complexed via charge interaction to introduce DNA into cells.

Also provided by this invention are conjugates that utilize the AAV5 capsid or a unique region of the AAV5 capsid protein (e.g. VP1, VP2 or VP3 or combinations thereof) to introduce DNA into cells. For example, the type 5 VP3 protein or fragment thereof, can be conjugated to a DNA on a plasmid that is conjugated to a lipid. Cells can be infected using the targeting ability of the VP3 capsid protein to achieve the desired tissue tropism, specific to AAV5. Type 5 VP1 and VP2 proteins can also be utilized to introduce DNA or other molecules into cells. By further incorporating the Rep protein and the AAV TRS into the DNA-containing conjugate, cells can be transduced and targeted integration can be achieved. For example, if AAV5 specific targeted integration is desired, a conjugate composed of the AAV5 VP3 capsid, AAV5 rep or a fragment of AAV5 rep, AAV5 TRS, the rep binding site, the heterologous DNA of interest, and a lipid, can be utilized to achieve AAV5 specific tropism and AAV5 specific targeted integration in the genome.

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Further provided by this invention are chimeric viruses where AAV5 can be combined with herpes virus, baculovirus or other viruses to achieve a desired tropism associated with another virus. For example, the AAV5 ITRs could be inserted in the herpes virus and cells could be infected. Post-infection, the ITRs of AAV5 could be acted on by AAV5 rep provided in the system or in a separate vehicle to rescue AAV5 from the genome. Therefore, the cellular tropism of the herpes simplex virus can be combined with AAV5 rep mediated targeted integration. Other viruses that could be

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utilized to construct chimeric viruses include, lentivirus, retrovirus, pseudotyped retroviral vectors, and adenoviral vectors.

The present invention further provides isolated nucleic acids of AAV5. For example, provided is an isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO:1 (AAV5 genome). This nucleic acid, or portions thereof, can be inserted into vectors, such as plasmids, yeast artificial chromosomes, or other viral vector (particle), if desired, by standard cloning methods. The present invention also provides an isolated nucleic acid consisting essentially of the nucleotide sequence set forth in SEQ ID NO:1. The nucleotides of SEQ ID NO:1 can have minor modifications and still be contemplated by the present invention. For example, modifications that do not alter the amino acid encoded by any given codon (such as by modification of the third, "wobble," position in a codon) can readily be made, and such alterations are known in the art. Furthermore, modifications that cause a resulting neutral (conserved) amino acid substitution of a similar amino acid can be made in a coding region of the genome. Additionally, modifications as described herein for the AAV5 components, such as the ITRs, the p5 promoter, etc. are contemplated in this invention. Furthermore, modifications to regions of SEQ ID NO:1 other than in the ITR, TRS Rep binding site and hairpin are likely to be tolerated without serious impact on the function of the nucleic acid as a recombinant vector.

As used herein, the term "isolated" refers to a nucleic acid separated or significantly free from at least some of the other components of the naturally occurring organism, for example, the cell structural components or viral components commonly found associated with nucleic acids in the environment of the virus and/or other nucleic acids. The isolation of the native nucleic acids can be accomplished, for example, by techniques such as cell lysis followed by phenol plus chloroform extraction, followed by ethanol precipitation of the nucleic acids. The nucleic acids of this invention can be isolated from cells according to any of many methods well known in the art.

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As used herein, the term "nucleic acid" refers to single-or multiple stranded molecules which may be DNA or RNA, or any combination thereof, including

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modifications to those nucleic acids. The nucleic acid may represent a coding strand or its complement, or any combination thereof. Nucleic acids may be identical in sequence to the sequences which are naturally occurring for any of the novel genes discussed herein or may include alternative codons which encode the same amino acid as those provided herein, including that which is found in the naturally occurring sequence. These nucleic acids can also be modified from their typical structure. Such modifications include, but are not limited to, methylated nucleic acids, the substitution of a non-bridging oxygen on the phosphate residue with either a sulfur (yielding phosphorothioate deoxynucleotides), selenium (yielding phosphorselenoate deoxynucleotides), or methyl groups (yielding methylphosphonate deoxynucleotides).

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The present invention additionally provides an isolated nucleic acid that selectively hybridizes with any nucleic acid disclosed herein, including the entire AAV5 genome and any unique fragment thereof, including the Rep and capsid encoding sequences (e.g. SEQ ID NOS: 1, 7, 8, 9, 10, 11, 13, 15, 16, 17, 18, 19, 20, 21, 15 22 and 23). Specifically, the nucleic acid can selectively or specifically hybridize to an isolated nucleic acid consisting of the nucleotide sequence set forth in SEQ ID NO:1 (AAV5 genome). The present invention further provides an isolated nucleic acid that selectively or specifically hybridizes with an isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO:1 (AAV5 genome). By "selectively 20 hybridizes" as used herein is meant a nucleic acid that hybridizes to one of the disclosed nucleic acids under sufficient stringency conditions without significant hybridization to a nucleic acid encoding an unrelated protein, and particularly, without detectably hybridizing to nucleic acids of AAV2. Thus, a nucleic acid that selectively hybridizes with a nucleic acid of the present invention will not selectively hybridize under stringent conditions with a nucleic acid encoding a different protein or the corresponding protein from a different serotype of the virus, and vice versa. A "specifically hybridizing" nucleic acid is one that hybridizes under stringent conditions to only a nucleic acid found in AAV5. Therefore, nucleic acids for use, for example, as primers and probes to detect or amplify the target nucleic acids are contemplated herein. Nucleic acid fragments that selectively hybridize to any given nucleic acid can be used, e.g., as primers and or probes for further hybridization or for amplification

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methods (e.g., polymerase chain reaction (PCR), ligase chain reaction (LCR)). Additionally, for example, a primer or probe can be designed that selectively hybridizes with both AAV5 and a gene of interest carried within the AAV5 vector (i.e., a chimeric nucleic acid).

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Stringency of hybridization is controlled by both temperature and salt concentration of either or both of the hybridization and washing steps. Typically, the stringency of hybridization to achieve selective hybridization involves hybridization in high ionic strength solution (6X SSC or 6X SSPE) at a temperature that is about 12-25°C below the T<sub>m</sub> (the melting temperature at which half of the molecules dissociate from their hybridization partners) followed by washing at a combination of temperature and salt concentration chosen so that the washing temperature is about 5°C to 20°C below the T<sub>m</sub>. The temperature and salt conditions are readily determined empirically in preliminary experiments in which samples of reference DNA immobilized on filters are hybridized to a labeled nucleic acid of interest and then washed under conditions of different stringencies. Hybridization temperatures are typically higher for DNA-RNA and RNA-RNA hybridizations. The washing temperatures can be used as described above to achieve selective stringency, as is known in the art. (Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989; Kunkel et al. Methods Enzymol. 1987:154:367, 1987). A preferable stringent hybridization condition for a DNA:DNA hybridization can be at about 68°C (in aqueous solution) in 6X SSC or 6X SSPE followed by washing at 68°C. Stringency of hybridization and washing, if desired, can be reduced accordingly as the degree of complementarity desired is decreased, and further, depending upon the G-C or A-T richness of any area wherein variability is searched for. Likewise, stringency of hybridization and washing, if desired, can be increased accordingly as homology desired is increased, and further, depending upon the G-C or A-T richness of any area wherein high homology is desired, all as known in the art.

A nucleic acid that selectively hybridizes to any portion of the AAV5 genome is contemplated herein. Therefore, a nucleic acid that selectively hybridizes to AAV5 can be of longer length than the AAV5 genome, it can be about the same length as the

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AAV5 genome or it can be shorter than the AAV5 genome. The length of the nucleic acid is limited on the shorter end of the size range only by its specificity for hybridization to AAV5, *i.e.*, once it is too short, typically less than about 5 to 7 nucleotides in length, it will no longer bind specifically to AAV5, but rather will hybridize to numerous background nucleic acids. Additionally contemplated by this invention is a nucleic acid that has a portion that specifically hybridizes to AAV5 and a portion that specifically hybridizes to a gene of interest inserted within AAV5.

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The present invention further provides an isolated nucleic acid encoding an adeno-associated virus 5 Rep protein. The AAV5 Rep proteins are encoded by open 10 reading frame (ORF) 1 of the AAV5 genome. Examples of the AAV5 Rep genes are shown in the nucleic acid set forth in SEQ ID NO:1, and include nucleic acids consisting essentially of the nucleotide sequences set forth in SEQ ID NOS:10 (Rep52). 11 (Rep78), 13 (Rep40), and 15 (Rep68), and nucleic acids comprising the nucleotide 15 sequences set forth in SEQ ID NOS:10, 11, 13, and 15. However, the present invention contemplates that the Rep nucleic acid can include any one, two, three, or four of the four Rep proteins, in any order, in such a nucleic acid. Furthermore, minor modifications are contemplated in the nucleic acid, such as silent mutations in the coding sequences, mutations that make neutral or conservative changes in the encoded 20 amino acid sequence, and mutations in regulatory regions that do not disrupt the expression of the gene. Examples of other minor modifications are known in the art. Further modifications can be made in the nucleic acid, such as to disrupt or alter expression of one or more of the Rep proteins in order to, for example, determine the effect of such a disruption; such as to mutate one or more of the Rep proteins to determine the resulting effect, etc. However, in general, a modified nucleic acid 25 encoding a Rep protein will have at least about 85%, about 90%, about 93%, about 95%, about 98% or 100% homology to the Rep nucleic sequences described herein e.g., SEQ ID NOS: 10, 11, 13 and 15, and the Rep polypeptide encoded therein will have overall about 93%, about 95%, about 98%, about 99% or 100% homology with the amino acid sequence described herein, e.g., SEQ ID NOS:2, 3, 12 and 14. Percent 30 homology is determined by the techniques described herein.

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The present invention also provides an isolated nucleic acid that selectively or specifically hybridizes with a nucleic acid consisting essentially of the nucleotide sequence set forth in SEQ ID NOS:10, 11, 13 and 15, and an isolated nucleic acid that selectively hybridizes with a nucleic acid comprising the nucleotide sequence set forth in SEQ ID NOS:10, 11, 13 and 15. "Selectively hybridizing" and "stringency of hybridization" is defined elsewhere herein.

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As described above, the present invention provides the nucleic acid encoding a Rep 40 protein and, in particular an isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO: 13, an isolated nucleic acid consisting essentially of the nucleotide sequence set forth in SEQ ID NO: 13, and a nucleic acid encoding the adeno-associated virus 5 protein having the amino acid sequence set forth in SEQ ID NO: 12. The present invention also provides the nucleic acid encoding a Rep 52 protein, and in particular an isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO:10, an isolated nucleic acid consisting essentially of the nucleotide sequence set forth in SEQ ID NO:10, and a nucleic acid encoding the adenoassociated virus 5 Rep protein having the amino acid sequence set forth in SEQ ID NO:2. The present invention further provides the nucleic acid encoding a Rep 68 protein and, in particular an isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO: 15, an isolated nucleic acid consisting essentially of the nucleotide sequence set forth in SEQ ID NO: 15, and a nucleic acid encoding the adeno-associated virus 5 protein having the amino acid sequence set forth in SEQ ID NO: 14. And, further, the present invention provides the nucleic acid encoding a Rep 78 protein, and in particular an isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO:11, an isolated nucleic acid consisting essentially of the nucleotide sequence set forth in SEQ ID NO:11, and a nucleic acid encoding the adeno-associated virus 5 Rep protein having the amino acid sequence set forth in SEQ ID NO:3. As described elsewhere herein, these nucleic acids can have minor modifications, including silent nucleotide substitutions, mutations causing conservative amino acid substitutions in the encoded proteins, and mutations in control regions that do not or minimally affect the encoded amino acid sequence.

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The present invention further provides a nucleic acid encoding the entire AAV5 Capsid polypeptide. Furthermore, the present invention provides a nucleic acid encoding each of the three AAV5 coat proteins, VP1, VP2, and VP3. Thus, the present invention provides a nucleic acid encoding AAV5 VP1, a nucleic acid encoding AAV5 VP2, and a nucleic acid encoding AAV5 VP3. Thus, the present invention provides a 5 nucleic acid encoding the amino acid sequence set forth in SEQ ID NO:4 (VP1); a nucleic acid encoding the amino acid sequence set forth in SEQ ID NO:5 (VP2), and a nucleic acid encoding the amino acid sequence set forth in SEQ ID NO:6 (VP3). The present invention also specifically provides a nucleic acid comprising SEQ ID NO:7 (VP1 gene); a nucleic acid comprising SEQ ID NO:8 (VP2 gene); and a nucleic acid 10 comprising SEQ ID NO:9 (VP3 gene). The present invention also specifically provides a nucleic acid consisting essentially of SEQ ID NO:7 (VP1 gene), a nucleic acid consisting essentially of SEQ ID NO:8 (VP2 gene), and a nucleic acid consisting essentially of SEQ ID NO:9 (VP3 gene). Minor modifications in the nucleotide sequences encoding the capsid, or coat, proteins are contemplated, as described above 15 for other AAV5 nucleic acids. However, in general, a modified nucleic acid encoding a capsid protein will have at least about 85%, about 90%, about 93%, about 95%, about 98% or 100% homology to the capsid nucleic sequences described herein e.g., SEQ ID NOS: 7, 8, and 9, and the capsid polypeptide encoded therein will have overall 20 about 93%, about 95%, about 98%, about 99% or 100% homology with the amino acid sequence described herein, e.g., SEQ ID NOS:4, 5, and 6. Nucleic acids that selectively hybridize with the nucleic acids of SEQ ID NOS:7,8 and 9 under the conditions described above are also provided.

The present invention also provides a cell containing one or more of the herein described nucleic acids, such as the AAV5 genome, AAV5 ORF1 and ORF2, each AAV5 Rep protein gene, or each AAV5 capsid protein gene. Such a cell can be any desired cell and can be selected based upon the use intended. For example, cells can include bacterial cells, yeast cells, insect cells, human HeLa cells and simian Cos cells as well as other human and mammalian cells and cell lines. Primary cultures as well as established cultures and cell lines can be used. Nucleic acids of the present invention can be delivered into cells by any selected means, in particular depending upon the

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target cells. Many delivery means are well-known in the art. For example, electroporation, calcium phosphate precipitation, microinjection, cationic or anionic liposomes, and liposomes in combination with a nuclear localization signal peptide for delivery to the nucleus can be utilized, as is known in the art. Additionally, if the nucleic acids are in a viral particle, the cells can simply be transduced with the virion by standard means known in the art for AAV transduction. Small amounts of the recombinant AAV5 virus can be made to infect cells and produce more of itself.

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The invention provides purified AAV5 polypeptides. The term "polypeptide" as used herein refers to a polymer of amino acids and includes full-length proteins and 10 fragments thereof. Thus, "protein," polypeptide," and "peptide" are often used interchangeably herein. Substitutions can be selected by known parameters to be neutral (see, e.g., Robinson WE Jr, and Mitchell WM., AIDS 4:S151-S162 (1990)). As will be appreciated by those skilled in the art, the invention also includes those polypeptides having slight variations in amino acid sequences or other properties. Such 15 variations may arise naturally as allelic variations (e.g., due to genetic polymorphism) or may be produced by human intervention (e.g., by mutagenesis of cloned DNA sequences), such as induced point, deletion, insertion and substitution mutants. Minor changes in amino acid sequence are generally preferred, such as conservative amino acid replacements, small internal deletions or insertions, and additions or deletions at 20 the ends of the molecules. Substitutions may be designed based on, for example, the model of Dayhoff, et al. (in Atlas of Protein Sequence and Structure 1978, Nat'l Biomed. Res. Found., Washington, D.C.). These modifications can result in changes in the amino acid sequence, provide silent mutations, modify a restriction site, or provide 25 other specific mutations. The location of any modifications to the polypeptide will often determine its impact on function. Particularly, alterations in regions non-essential to protein function will be tolerated with fewer effects on function. Elsewhere in the application regions of the AAV5 proteins are described to provide guidance as to where substitutions, additions or deletions can be made to minimize the likelihood of 30 disturbing the function of the variant.

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A polypeptide of the present invention can be readily obtained by any of several means. For example, the polypeptide of interest can be synthesized chemically by standard methods. Additionally, the coding regions of the genes can be recombinantly expressed and the resulting polypeptide isolated by standard methods. Furthermore, an antibody specific for the resulting polypeptide can be raised by standard methods (see, e.g., Harlow and Lane, Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1988), and the protein can be isolated from a cell expressing the nucleic acid encoding the polypeptide by selective hybridization with the antibody. This protein can be purified to the extent desired by standard methods of protein purification (see, e.g., Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor Laboratory, Cold Spring Harbor, New York, 1989).

Typically, to be unique, a polypeptide fragment of the present invention will be
at least about 5 amino acids in length; however, unique fragments can be 6, 7, 8, 9, 10,
20, 30, 40, 50, 60, 70, 80, 90, 100 or more amino acids in length. A unique polypeptide
will typically comprise such a unique fragment; however, a unique polypeptide can also
be determined by its overall homology. A unique polypeptide can be 6, 7, 8, 9, 10, 20,
30, 40, 50, 60, 70, 80, 90, 100 or more amino acids in length. Uniqueness of a
polypeptide fragment can readily be determined by standard methods such as searches
of computer databases of known peptide or nucleic acid sequences or by hybridization
studies to the nucleic acid encoding the protein or to the protein itself, as known in the
art. The uniqueness of a polypeptide fragment can also be determined immunologically
as well as functionally. Uniqueness can be simply determined in an amino acid-byamino acid comparison of the polypeptides.

An antigenic or immunoreactive fragment of this invention is typically an amino acid sequence of at least about 5 consecutive amino acids, and it can be derived from the AAV5 polypeptide amino acid sequence. An antigenic AAV5 fragment is any fragment unique to the AAV5 protein, as described herein, against which an AAV5-specific antibody can be raised, by standard methods. Thus, the resulting antibody-antigen reaction should be specific for AAV5.

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The present invention provides an isolated AAV5 Rep protein. An AAV5 Rep polypeptide is encoded by ORF1 of AAV5. The present invention also provides each individual AAV5 Rep protein. Thus the present invention provides AAV5 Rep 40 (e.g., SEQ ID NO: 12), or a unique fragment thereof. The present invention provides AAV5 Rep 52 (e.g., SEQ ID NO: 2), or a unique fragment thereof. The present invention provides AAV5 Rep 68 (e.g., SEQ ID NO: 14), or a unique fragment thereof. The present invention provides an example of AAV5 Rep 78 (e.g., SEQ ID NO: 3), or a unique fragment thereof. By "unique fragment thereof" is meant any smaller polypeptide fragment encoded by an AAV5 rep gene that is of sufficient length to be found only in the Rep polypeptide. Substitutions and modifications of the amino acid sequence can be made as described above and, further, can include protein processing modifications, such as glycosylation, to the polypeptide.

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The present invention further provides an AAV5 Capsid polypeptide or a unique fragment thereof. AAV5 capsid polypeptide is encoded by ORF 2 of AAV5. 15 The present invention further provides the individual AAV5 capsid proteins, VP1, VP2 and VP3 or unique fragments thereof. Thus, the present invention provides an isolated polypeptide having the amino acid sequence set forth in SEQ ID NO:4 (VP1). The present invention additionally provides an isolated polypeptide having the amino acid 20 sequence set forth in SEQ ID NO:5 (VP2). The present invention also provides an isolated polypeptide having the amino acid sequence set forth in SEQ ID NO:6 (VP3). By "unique fragment thereof" is meant any smaller polypeptide fragment encoded by any AAV5 capsid gene that is of sufficient length to be found only in the AAV5 capsid protein. Substitutions and modifications of the amino acid sequence can be made as described above and, further, can include protein processing modifications, such as 25 glycosylation, to the polypeptide. However, an AAV5 Capsid polypeptide including all three coat proteins will have greater than about 56% overall homology to the polypeptide encoded by the nucleotides set forth in SEQ ID NOS:4,5 or 6. The protein can have about 65%, about 70%, about 75%, about 80%, about 85%, about 90%, 93%, 95%, 97% or even 100% homology to the amino acid sequence encoded by the 30 nucleotides set forth in SEQ ID NOS:4,5 or 6. An AAV5 VP1 polypeptide can have at least about 58%, about 60%, about 70%, about 80%, about 90%, 93%, 95%, 97% or

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about 100% homology to the amino acid sequence set forth in SEQ ID NO:4. An AAV5 VP2 polypeptide can have at least about 58%, about 60%, about 70%, about 80%, about 90%, 93%, 95%, 97% or about 100% homology to the amino acid sequence set forth in SEQ ID NO:5. An AAV5 VP3 polypeptide can have at least about 60%, about 70%, about 80%, about 90%, 93%, 95%, 97% or about 100% homology to the amino acid sequence set forth in SEQ ID NO:6.

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The present invention further provides an isolated antibody that specifically binds an AAV5 Rep protein or a unique epitope thereof. Also provided are isolated antibodies that specifically bind the AAV5 Rep 52 protein, the AAV5 Rep 40 protein, the AAV5 Rep 68 protein and the AAV5 Rep 78 protein having the amino acid sequences set forth in SEQ ID NO: 2, SEQ ID NO: 12, SEQ ID NO: 14 and SEQ ID NO: 3, respectively or that specifically binds a unique fragment thereof. Clearly, any given antibody can recognize and bind one of a number of possible epitopes present in the polypeptide; thus only a unique portion of a polypeptide (having the epitope) may need to be present in an assay to determine if the antibody specifically binds the polypeptide.

The present invention additionally provides an isolated antibody that specifically binds any of the adeno-associated virus 5 Capsid proteins (VP1, VP2 or 20 VP3), a unique epitope thereof, or the polypeptide comprising all three AAV5 coat proteins. Also provided is an isolated antibody that specifically binds the AAV5 capsid protein having the amino acid sequence set forth in SEQ ID NO:4 (VP1), or that specifically binds a unique fragment thereof. The present invention further provides an isolated antibody that specifically binds the AAV5 Capsid protein having the amino 25 acid sequence set forth in SEQ ID NO:5 (VP2), or that specifically binds a unique fragment thereof. The invention additionally provides an isolated antibody that specifically binds the AAV5 Capsid protein having the amino acid sequence set forth in SEQ ID NO:6 (VP3), or that specifically binds a unique fragment thereof. Again, any given antibody can recognize and bind one of a number of possible epitopes present in 30 the polypeptide; thus only a unique portion of a polypeptide (having the epitope) may

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need to be present in an assay to determine if the antibody specifically binds the polypeptide.

The antibody can be a component of a composition that comprises an antibody that specifically binds the AAV5 protein. The composition can further comprise, *e.g.*, serum, serum-free medium, or a pharmaceutically acceptable carrier such as physiological saline, etc..

By "an antibody that specifically binds" an AAV5 polypeptide or protein is meant an antibody that selectively binds to an epitope on any portion of the AAV5 10 peptide such that the antibody binds specifically to the corresponding AAV5 polypeptide without significant background. Specific binding by an antibody further means that the antibody can be used to selectively remove the target polypeptide from a sample comprising the polypeptide or and can readily be determined by radioimmunoassay (RIA), bioassay, or enzyme-linked immunosorbant (ELISA) 15 technology. An ELISA method effective for the detection of the specific antibodyantigen binding can, for example, be as follows: (1) bind the antibody to a substrate; (2) contact the bound antibody with a sample containing the antigen; (3) contact the above with a secondary antibody bound to a detectable moiety (e.g., horseradish 20 peroxidase enzyme or alkaline phosphatase enzyme); (4) contact the above with the substrate for the enzyme; (5) contact the above with a color reagent; (6) observe the color change.

An antibody can include antibody fragments such as Fab fragments which retain
the binding activity. Antibodies can be made as described in, e.g., Harlow and Lane,
Antibodies: A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring
Harbor, New York (1988). Briefly, purified antigen can be injected into an animal in
an amount and in intervals sufficient to elicit an immune response. Antibodies can
either be purified directly, or spleen cells can be obtained from the animal. The cells
are then fused with an immortal cell line and screened for antibody secretion.
Individual hybridomas are then propagated as individual clones serving as a source for
a particular monoclonal antibody.

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The present invention additionally provides a method of screening a cell for infectivity by AAV5 comprising contacting the cell with AAV5 and detecting the presence of AAV5 in the cells. AAV5 particles can be detected using any standard physical or biochemical methods. For example, physical methods that can be used for this detection include DNA based methods such as 1) polymerase chain reaction (PCR) for viral DNA or RNA or 2) direct hybridization with labeled probes, and immunological methods such as by 3) antibody directed against the viral structural or non- structural proteins. Catalytic methods of viral detection include, but are not limited to, detection of site and strand specific DNA nicking activity of Rep proteins or replication of an AAV origin- containing substrate. Reporter genes can also be utilized to detect cells that transduct AAV-5. For example, β-gal, green flourescent protein or luciferase can be inserted into a recombinant AAV-5. The cell can then be contacted with the recombinant AAV-5, either in vitro or in vivo and a colorimetric assay could detect a color change in the cells that would indicate transduction of AAV-5 in the cell. Additional detection methods are outlined in Fields, Virology, Raven Press, New York, New York. 1996.

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For screening a cell for infectivity by AAV5, wherein the presence of AAV5 in the cells is determined by nucleic acid hybridization methods, a nucleic acid probe for such detection can comprise, for example, a unique fragment of any of the AAV5 nucleic acids provided herein. The uniqueness of any nucleic acid probe can readily be determined as described herein. Additionally, the presence of AAV5 in cells can be determined by flourescence, antibodies to gene products, focus forming assays, plaque lifts, Western blots and chromogenic assays. The nucleic acid can be, for example, the nucleic acid whose nucleotide sequence is set forth in SEQ ID NO: 1,7, 8, 9, 10, 11, 13, 15, 16, 17, 18, 19, 20, 21, 22, 23 or a unique fragment thereof.

The present invention includes a method of determining the suitability of an AAV5 vector for administration to a subject comprising administering to an antibody-containing sample from the subject an antigenic fragment of an isolated AAV5 Rep or Capsid protein, and detecting neutralizing antibody-antigen reaction in the sample, the presence of a neutralizing reaction indicating the AAV5 vector may be unsuitable for

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use in the subject. The present method of determining the suitability of an AAV5 vector for administration to a subject can comprise contacting an antibody-containing sample from the subject with a unique antigenic or immunogenic fragment of an AAV5 Rep protein (e.g. Rep 40, Rep 52, Rep 68, Rep 78) and detecting an antibody-antigen reaction in the sample, the presence of a reaction indicating the AAV5 vector to be unsuitable for use in the subject. The AAV5 Rep proteins are provided herein, and their antigenic fragments are routinely determined. The AAV5 capsid protein can be used to select an antigenic or immunogenic fragment, for example from the amino acid sequence set forth in SEQ ID NO:4 (VP1), the amino acid sequence set forth in SEQ ID NO: 5 (VP2) or the amino acid sequence set forth in SEQ ID NO:6 (VP3). 10 Alternatively, or additionally, an antigenic or immunogenic fragment of an isolated AAV5 Rep protein can be utilized in this determination method. The AAV5 Rep protein from which an antigenic fragment is selected can have the amino acid sequence encoded by the nucleic acid set forth in SEQ ID NO:1, the amino acid sequence set 15 forth in SEQ ID NO:2, or the amino acid sequence set forth in SEQ ID NO:3, the amino acid sequence set forth in SEQ ID NO: 12, or the amino acid sequence set forth in SEQ ID NO:14.

antigenicity, immunogenicity and/or specificity. Briefly, various concentrations of a putative immunogenically specific fragment are prepared and administered to a subject and the immunological response (e.g., the production of antibodies or cell mediated immunity) of an animal to each concentration is determined. The amounts of antigen administered depend on the subject, e.g. a human, rabbit or a guinea pig, the condition of the subject, the size of the subject, etc. Thereafter an animal so inoculated with the antigen can be exposed to the AAV5 viral particle or AAV5 protein to test the immunoreactivity or the antigenicity of the specific immunogenic fragment. The specificity of a putative antigenic or immunogenic fragment can be ascertained by testing sera, other fluids or lymphocytes from the inoculated animal for cross reactivity with other closely related viruses, such as AAV1, AAV2, AAV3, AAV4 and AAV5.

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The hemagglutination assay can also be used to rapidly identify and detect AAV5 viral particles. Detection of hemagglutination activity correlates with infectivity and can be used to titer the virus. This assay could also be used to identify antibodies in a patients serum which might interact with the virus. Hemagglutination has been shown to correlate with infectivity and therefore hemagglutination maybe a useful assay for identify cellular receptors for AAV5.

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By the "suitability of an AAV5 vector for administration to a subject" is meant a determination of whether the AAV5 vector will elicit a neutralizing immune response upon administration to a particular subject. A vector that does not elicit a significant immune response is a potentially suitable vector, whereas a vector that elicits a significant, neutralizing immune response (e.g. at least 90%) is thus likely to be unsuitable for use in that subject. Significance of any detectable immune response is a standard parameter understood by the skilled artisan in the field. For example, one can incubate the subject's serum with the virus, then determine whether that virus retains its ability to transduce cells in culture. If such virus cannot transduce cells in culture, the vector likely has elicited a significant immune response.

Alternatively, or additionally, one skilled in the art could determine whether or not AAV5 administration would be suitable for a particular cell type of a subject. For example, the artisan could culture muscle cells *in vitro* and transduce the cells with AAV5 in the presence or absence of the subject's serum. If there is a reduction in transduction efficiency, this could indicate the presence of a neutralizing antibody or other factors that may inhibit transduction. Normally, greater than 90% inhibition would have to be observed in order to rule out the use of AAV-5 as a vector. However, this limitation could be overcome by treating the subject with an immunosuppressant that could block the factors inhibiting transduction.

As will be recognized by those skilled in the art, numerous types of
immunoassays are available for use in the present invention to detect binding between
an antibody and an AAV5 polypeptide of this invention. For instance, direct and
indirect binding assays, competitive assays, sandwich assays, and the like, as are

generally described in, *e.g.*, U.S. Pat. Nos. 4,642,285; 4,376,110; 4,016,043; 3,879,262; 3,852,157; 3,850,752; 3,839,153; 3,791,932; and Harlow and Lane, *Antibodies, A Laboratory Manual*, Cold Spring Harbor Publications, N.Y. (1988). For example, enzyme immunoassays such as immunofluorescence assays (IFA), enzyme linked immunosorbent assays (ELISA) and immunoblotting can be readily adapted to accomplish the detection of the antibody. An ELISA method effective for the detection of the antibody bound to the antigen can, for example, be as follows: (1) bind the antigen to a substrate; (2) contact the bound antigen with a fluid or tissue sample containing the antibody; (3) contact the above with a secondary antibody specific for the antigen and bound to a detectable moiety (e.g., horseradish peroxidase enzyme or alkaline phosphatase enzyme); (4) contact the above with the substrate for the enzyme; (5) contact the above with a color reagent; (6) observe color change.

The antibody-containing sample of this method can comprise any biological sample which would contain the antibody or a cell containing the antibody, such as blood, plasma, serum, bone marrow, saliva and urine.

The present invention also provides a method of producing the AAV5 virus by transducing a cell with the nucleic acid encoding the virus.

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The present method further provides a method of delivering an exogenous (heterologous) nucleic acid to a cell comprising administering to the cell an AAV5 particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to the cell.

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The AAV ITRs in the vector for the herein described delivery methods can be AAV5 ITRs (SEQ ID NOS: 19 and 20). Furthermore, the AAV ITRs in the vector for the herein described nucleic acid delivery methods can also comprise AAV1, AAV2, AAV3, AAV4, or AAV6 inverted terminal repeats.

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The present invention also includes a method of delivering a heterologous nucleic acid to a subject comprising administering to a cell from the subject an AAV5

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particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, and returning the cell to the subject, thereby delivering the nucleic acid to the subject. The AAV ITRs can be any AAV ITRs, including AAV5 ITRs and AAV2 ITRs. For example, in an ex vivo administration, cells are isolated from a subject by standard means according to the cell type and placed in appropriate 5 culture medium, again according to cell type (see, e.g., ATCC catalog). Viral particles are then contacted with the cells as described above, and the virus is allowed to transduce the cells. Cells can then be transplanted back into the subject's body, again by means standard for the cell type and tissue (e. g., in general, U.S. Patent No. 5,399,346; for neural cells, Dunnett, S.B. and Björklund, A., eds., Transplantation: 10 Neural Transplantation-A Practical Approach, Oxford University Press, Oxford (1992)). If desired, prior to transplantation, the cells can be studied for degree of transduction by the virus, by known detection means and as described herein. Cells for ex vivo transduction followed by transplantation into a subject can be selected from those listed above, or can be any other selected cell. Preferably, a selected cell type is 15 examined for its capability to be transfected by AAV5. Preferably, the selected cell will be a cell readily transduced with AAV5 particles; however, depending upon the application, even cells with relatively low transduction efficiencies can be useful, particularly if the cell is from a tissue or organ in which even production of a small amount of the protein or antisense RNA encoded by the vector will be beneficial to the 20 subject.

The present invention further provides a method of delivering a nucleic acid to a cell in a subject comprising administering to the subject an AAV5 particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to a cell in the subject. Administration can be an *ex vivo* administration directly to a cell removed from a subject, such as any of the cells listed above, followed by replacement of the cell back into the subject, or administration can be *in vivo* administration to a cell in the subject. For *ex vivo* administration, cells are isolated from a subject by standard means according to the cell type and placed in appropriate culture medium, again according to cell type (*see*, *e.g.*, ATCC catalog). Viral particles are then contacted with the cells as described above,

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and the virus is allowed to transfect the cells. Cells can then be transplanted back into the subject's body, again by means standard for the cell type and tissue (e. g., for neural cells, Dunnett, S.B. and Björklund, A., eds., *Transplantation: Neural Transplantation-A Practical Approach*, Oxford University Press, Oxford (1992)). If desired, prior to transplantation, the cells can be studied for degree of transfection by the virus, by known detection means and as described herein.

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The present invention further provides a method of delivering a nucleic acid to a cell in a subject having neutralizing antibodies to AAV2 comprising administering to the subject an AAV5 particle containing a vector comprising the nucleic acid, thereby 10 delivering the nucleic acid to a cell in the subject. A subject that has neutralizing antibodies to AAV2 can readily be determined by any of several known means, such as contacting AAV2 protein(s) with an antibody-containing sample, such as blood, from a subject and detecting an antigen-antibody reaction in the sample. Delivery of the AAV5 particle can be by either ex vivo or in vivo administration as herein described. 15 Thus, a subject who might have an adverse immunogenic reaction to a vector administered in an AAV2 viral particle can have a desired nucleic acid delivered using an AAV5 particle. This delivery system can be particularly useful for subjects who have received therapy utilizing AAV2 particles in the past and have developed antibodies to AAV2. An AAV5 regimen can now be substituted to deliver the desired 20 nucleic acid.

In any of the methods of delivering heterologous nucleic acids to a cell or subject described herein, the AAV5-conjugated nucleic acid or AAV5 particle-conjugated nucleic acids described herein can be used.

In vivo administration to a human subject or an animal model can be by any of many standard means for administering viruses, depending upon the target organ, tissue or cell. Virus particles can be administered orally, parenterally (e.g., intravenously), by intramuscular injection, by direct tissue or organ injection, by intraperitoneal injection, topically, transdermally, via aerosol delivery, via the mucosa or the like. Viral nucleic acids (non-encapsidated) can also be administered, e.g., as a complex with cationic

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liposomes, or encapsulated in anionic liposomes. The present compositions can include various amounts of the selected viral particle or non-encapsidated viral nucleic acid in combination with a pharmaceutically acceptable carrier and, in addition, if desired, may include other medicinal agents, pharmaceutical agents, carriers, adjuvants, diluents, etc.

5 Parental administration, if used, is generally characterized by injection. Injectables can be prepared in conventional forms, either as liquid solutions or suspensions, solid forms suitable for solution or suspension in liquid prior to injection, or as emulsions. Dosages will depend upon the mode of administration, the disease or condition to be treated, and the individual subject's condition, but will be that dosage typical for and used in administration of other AAV vectors, such as AAV2 vectors. Often a single dose can be sufficient; however, the dose can be repeated if desirable.

Administration methods can be used to treat brain disorders such as Parkinson's disease, Alzheimer's disease, and demyelination disease. Other diseases that can be treated by these methods include metabolic disorders such as , muscoloskeletal diseases, cardiovascular disease, cancer, and autoimmune disorders.

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Administration of this recombinant AAV5 virion to the cell can be accomplished by any means, including simply contacting the particle, optionally contained in a desired liquid such as tissue culture medium, or a buffered saline solution, with the cells. The virion can be allowed to remain in contact with the cells for any desired length of time, and typically the virion is administered and allowed to remain indefinitely. For such *in vitro* methods, the virion can be administered to the cell by standard viral transduction methods, as known in the art and as exemplified herein. Titers of virus to administer can vary, particularly depending upon the cell type, but will be typical of that used for AAV transduction in general which is well known in the art. Additionally the titers used to transduce the particular cells in the present examples can be utilized.

The cells that can be transduced by the present recombinant AAV5 virion can include any desired cell, such as the following cells and cells derived from the following tissues, human as well as other mammalian tissues, such as primate, horse,

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sheep, goat, pig, dog, rat, and mouse: Adipocytes, Adenocyte, Adrenal cortex, Amnion, Aorta, Ascites, Astrocyte, Bladder, Bone, Bone marrow, Brain, Breast, Bronchus, Cardiac muscle, Cecum, Cervix, Chorion, Colon, Conjunctiva, Connective tissue, Cornea, Dermis, Duodenum, Endometrium, Endothelial cells, Epithelial tissue, Epithelial cells, Epidermis, Esophagus, Eye, Fascia, Fibroblasts, Foreskin, 5 Gastric, Glial cells, Glioblast, Gonad, Hepatic cells, Histocyte, Ileum, Intestine, small Intestine, Jejunum, Keratinocytes, Kidney, Larynx, Leukocytes, Lipocyte, Liver, Lung, Lymph node, Lymphoblast, Lymphocytes, Macrophages, Mammary alveolar nodule, Mammary gland, Mastocyte, Maxilla, Melanocytes, Mesenchymal, Monocytes, Mouth, 10 Myelin, Myoblasts Nervous tissue, Neuroblast, Neurons, Neuroglia, Osteoblasts, Osteogenic cells, Ovary, Palate, Pancreas, Papilloma, Peritoneum, Pituicytes, Pharynx, Placenta, Plasma cells, Pleura, Prostate, Rectum, Salivary gland, Skeletal muscle, Skin, Smooth muscle, Somatic, Spleen, Squamous, Stomach, Submandibular gland, Submaxillary gland, Synoviocytes, Testis, Thymus, Thyroid, Trabeculae, Trachea, 15 Turbinate, Umbilical cord, Ureter, and Uterus.

#### STATEMENT OF UTILITY

The present invention provides recombinant vectors based on AAV5. Such 20 vectors may be useful for transducing erythroid progenitor cells or cells lacking heparin sulfate proteoglycans which is very inefficient with AAV2 based vectors. These vectors may also be useful for transducing cells with a nucleic acid of interest in order to produce cell lines that could be used to screen for agents that interact with the gene product of the nucleic acid of interest. In addition to transduction of other cell types, 25 transduction of erythroid cells would be useful for the treatment of cancer and genetic diseases which can be corrected by bone marrow transplants using matched donors. Some examples of this type of treatment include, but are not limited to, the introduction of a therapeutic gene such as genes encoding interferons, interleukins, tumor necrosis factors, adenosine deaminase, cellular growth factors such as lymphokines, blood coagulation factors such as factor VIII and IX, cholesterol metabolism uptake and 30 transport protein such as EpoE and LDL receptor, and antisense sequences to inhibit viral replication of, for example, hepatitis or HIV.

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The present invention provides a vector comprising the AAV5 virus as well as AAV5 viral particles. While AAV5 is similar to AAV2, the two viruses are found herein to be physically and genetically distinct. These differences endow AAV5 with some unique advantages which better suit it as a vector for gene therapy.

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Furthermore, as shown herein, AAV5 capsid protein is distinct from AAV2 capsid protein and exhibits different tissue tropism. AAV2 and AAV5 likely utilize distinct cellular receptors. AAV2 and AAV5 are serologically distinct and thus, in a gene therapy application, AAV5 would allow for transduction of a patient who already possess neutralizing antibodies to AAV2 either as a result of natural immunological defense or from prior exposure to AAV2 vectors.

The present invention is more particularly described in the following examples which are intended as illustrative only since numerous modifications and variations therein will be apparent to those skilled in the art.

#### **EXAMPLES**

To understand the nature of AAV5 virus and to determine its usefulness as a vector for gene transfer, it was cloned and sequenced.

Cell culture and virus propagation

Cos and HeLa cells were maintained as monolayer cultures in D10 medium (Dulbecco's modified Eagle's medium containing 10% fetal calf serum, 100  $\mu$ g/ml penicillin, 100 units/ml streptomycin and IX Fungizone as recommended by the manufacturer; (GIBCO, Gaithersburg, MD, USA) . All other cell types were grown under standard conditions which have been previously reported.

Virus was produced as previously described for AAV2 using the Beta 30 galactosidase vector plasmid and a helper plasmid containing the AAV5 Rep and Cap genes (9). The helper plasmid was constructed in such a way to minimize any homologous sequence between the helper and vector plasmids. This step was taken to WO 99/61601

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minimize the potential for wild-type (wt) particle formation by homologous recombination.

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## DNA Cloning and Sequencing and Analysis

In order to clone the genome of AAV5, infectious cell lysate was expanded in adherent cos cells and then suspension HeLa cells with the resulting viral particles isolated by CsCl isopynic gradient centrifugation. DNA dot blots of Aliquots of the gradient fractions indicated that the highest concentration of viral genomes were contained in fractions with a refractive index of approx. 1.372. While the initial description of the virus did not determine the density of the particles, this value is similar to that of AAV2. Analysis of annealed virion derived DNA obtained from these fractions indicated a major species of 4.6 kb in length which upon restriction analysis gave bands similar in size to those previously reported. Additional restriction mapping indicated a unique BssHII site at one end of the viral genome. This site was used to clone the major fragment of the viral genome. Additional overlapping clones were isolated and the sequence determined. Two distinct open reading frames (ORF) were identified. Computer analysis indicated that the left-hand ORF is approx 60% similar to that of the Rep gene of AAV2. Of the 4 other reported AAV serotypes, all have greater than 90% similarity in this ORF. The right ORF of the viral capsid proteins is also approximately 60% homologous to the Capsid ORF of AAV2. As with other AAV serotypes reported, the divergence between AAV5 and AAV2 is clustered in multiple blocks. By using the published three dimensional structure of the canine parvovirus and computer aided sequence comparisons, a number of these divergent regions have been shown to be on the exterior of the virus and thus suggest an altered tissue tropism.

Within the p5 promoter, a number of the core transcriptional elements are conserved such as the tataa box and YY1 site around the transcriptional start site. However the YY1 site at -60 and the upstream E-Box elements are not detectable suggesting an alternative method of regulation or activation.

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The inverted terminal repeats (ITRs) of the virus were cloned as a fragment from the right end of the genome. The resulting fragment was found to contain a number of sequence changes compared to AAV2. However, these changes were found to be complementary and did not affect the ability of this region to fold into a hairpin structure. Within the stem region of the hairpin two sequence elements have been found to be critical for the function of the ITRs as origins of viral replication. A repeat motif of GAGC/T which serves as the recognition site of Rep and a GGTTGAG sequence downstream of the Rep binding site which is the position of Rep's site and strand specific cleavage reaction. This sequence is not conserved between AAV5 and the other cloned AAV's suggesting that the ITRs and Rep proteins of AAV5 cannot compliment the other known AAV's.

To test the cross complementarity of AAV2 ITR containing genome and AAV5 ITR containing genomes recombinant particles were packaged either using type 2 Rep and Cap or type 5 Rep and Cap expression plasmids as previously described. As shown in Fig. 2, viral particles were produced only when the respective expression plasmids were used to package the cognate ITRs. This result is distinct from that of other serotypes of AAV which have shown cross complementary in packaging.

This specificity of AAV5 Rep for AAV5 ITRs was confirmed using a terminal resolution assay which can identify the site within one ITR cleaved by the Rep protein. Incubation of the Type 5 Rep protein with a type 2 ITR did not produce any cleavage products. In contrast, addition of type 2 Rep cleaved the DNA at the expected site. However AAV5 Rep did produce cleavage products when incubated with a type 5 ITR.

The site mapped to a region 21 bases from the Rep binding motif that is similar to AAV2 TRS. The site in AAV2 is CGGT TGAG (SEQ ID NO: 22) but in type 5 ITR is CGGT GTGA (SEQ ID NO: 21). The ability of AAV5 Rep to cleave at a different but similarly positioned site may result in integration of AAV5 at a distinct chromosomal locus compared to AAV2.

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Recombinant virus produced using AAV5 Rep and Cap was obtained at a greater titer than type 2. For example, in a comparative study, virus was isolated from

8X10<sup>7</sup> COS cells by CsCl banding and the distribution of the Beta galactosidase genomes across the gradient were determined by DNA dot blots of aliquots of gradient fractions. DNA dot blot titers indicated that AAV5 particles were produced at a 10-50 fold higher level than AAV2.

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The sequence divergence in the capsid protein ORF implies that the tissue tropism of AAV2 and AAV5 would differ. To study the transduction efficiency of AAV5 and AAV2, a variety of cell lines were transduced with serial dilution's of the purified virus expressing the gene for nuclear localized Beta galactosidase activity. Approx. 2X10<sup>4</sup> cells were exposed to virus in 1 ml of serum containing media for a 10 period of 48-60 hrs. After this time the cells were fixed and stained for Beta-galactosidase activity with 5-Bromo-4-chloro- 3-indolyl-b-D- galactopyranoside (Xgal) (ICN Biochemicals). Biological titers were determined by counting the number of positive cells in the different dilutions using a calibrated microscope ocular then multiplying by the area of the well. Titers were determined by the average number of cells in a minimum of 10 fields/well. Transduction of cos, HeLa, and 293, and IB3 cells with a similar number of particles showed approximately 10 fold decrease in titer with AAV5 compared with AAV2. In contrast MCF7 cells showed a 50-100 fold difference in transduction efficiency. Furthermore, both vectors transduced NIH 3T3 cells relatively poorly.

A recent publication reported that heparin proteoglycans on the surface of cells are involved in viral transduction. Addition of soluble heparin has been shown to inhibit transduction by blocking viral binding. Since the transduction data suggested a difference in tissue tropism for AAV5 and AAV2, the sensitivity of AAV5 transduction to heparin was determined. At an MOI of 100, the addition of 20µg/ml of heparin had no effect on AAV5 transduction. In contrast this amount of heparin inhibited 90% of the AAV2 transduction. Even at an MOI of 1000, no inhibition of AAV5 transduction was detected. These data support the conclusions of the tissue tropism study, i.e. that AAV2 and AAV5 may utilize a distinct cell surface molecules and therefore the mechanism of uptake may differ as well.

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AAV5 is a distinct virus within the dependovirus family based on sequence analysis, tissue tropism, and sensitivity to heparin. While elements of the P5 promoter are retained between AAV2-6 some elements are absent in AAV5 suggesting alternative mechanism of regulation. The ITR and Rep ORF are distinct from those previously identified and fail to complement the packaging of AAV2 based genomes. The ITR of AAV5 contains a different TRS compared to other serotypes of AAV which is responsible for the lack of complementation of the ITRs. This unique TRS should also result in a different integration locus for AAV5 compared to that of AAV2. Furthermore the production of recombinant AAV5 particles using standard packaging systems is approx. 10-50 fold better than AAV2. The majority of the differences in the capsid proteins lies in regions which have been proposed to be on the exterior of the surface of the parvovirus. These changes are most likely responsible for the lack of cross reactive antibodies and altered tissue tropism compared to AAV2.

ID NO: 18) produces rep 68 (a spliced site mutant) and rep78 and the p19 promoter (SEQ ID NO: 16) produces rep 40 (a spliced site mutant) and rep 52. While these regions are not well conserved within the Rep ORF of AAV5 some splice acceptor and donor sites exist in approximately the same region as the AAV2 sites. These sites can be identified using standard computer analysis programs such as signal in the PCGENE program. Therefore the sequences of the Rep proteins can be routinely identified as in other AAV serotypes.

#### Hemagglutination assay

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Hemagglutination activity was measured essentially as described previously (Chiorini et al 1997 J. Virol. Vol 71 6823-6833) Briefly 2 fold serial dilutions of virus in EDTA-buffered saline were mixed with an equal volume of 0.4% red blood cells in plastic U-bottom 96 well plates. The reaction was complete after a 2-h incubation at 8°C. Addition of purified AAV5 to a hemagglutination assay resulted in hemagglutination activity.

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## Transduction of airway epithelial cells

Primary airway epithilial cells were cultured and plated as previously described (Fasbender et al. J. Clin Invest. 1998 Jul 1; 102 (1): 184-93). Cells were transducted with an equivalent number of rAAV2 or rAAV5 particles containing a nuclear localized  $\beta$ -gal transgene with 50 particles of virus/cell (MOI 50) and continued in culture for 10 days.  $\beta$ -gal activity was determined following the procedure of (Chiorini et al. 1995 HGT Vol: 6 1531-1541) and the relative transduction efficiency compared. As shown in Figure 7, AAV5 transduced these cells 50- fold more efficiently than AAV2. This is the first time apical cells or cells exposed to the air have been shown to be infected by a gene therapy agent.

## Transduction of striated muscle

Chicken myoblasts were cultured and plated as previously described (Rhodes & Yamada 1995 NAR Vol 23 (12) 2305-13). Cells were allowed to fuse and then transduced with a similar number of particles of rAAV2 or rAAV5 containing a nuclear localized  $\beta$ -gal transgene as previously described above after 5 days in culture. The cells were stained for  $\beta$ -gal activity following the procedure of (Chiorini et al. 1995 HGT Vol: 6 1531-1541) and the relative transduction efficiency compared. As shown in Figure 8, AAV5 transduced these cells approximately 16 fold more efficiently than AAV2.

#### Transduction of rat brain explants

Primary neonatal rat brain explants were prepared as previously described (Scortegagna et al. Neurotoxicology. 1997; 18 (2): 331-9). After 7 days in culture, cells were transduced with a similar number of particles of rAAV5 containing a nuclear localized  $\beta$ -gal transgene as previously described. After 5 days in culture, the cells were stained for  $\beta$ -gal activity following the procedure of (Chiorini et al. 1995 HGT Vol: 6 1531-1541). As shown in Figure 9, transduction was detected in a variety of cell types including astrocytes, neuronal cels and glial cells.

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Transduction of human umbilical vein endothelial cells

Human umbilical vein endothelial cells were cultured and plated as previously described (Gnantenko et al. J Investig Med. 1997 Feb; 45(2): 87-98). Cells were transduced with rAAV2 or rAAV5 containing a nuclear localized  $\beta$ -gal transgene with 10 particles of virus/ cell (MOI 5) in minimal media then returned to complete media. After 24 hrs in culture the cells were stained for  $\beta$ -gal activity following the procedure of Chiorini et al. (1995 HGT Vol: 6 1531-1541), and the relative transduction efficiency compared. As shown in Figure 10, AAV5 transduced these cell 5-10 fold more efficiently than AAV2.

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Throughout this application, various publications are referenced. The disclosures of these publications in their entireties are hereby incorporated by reference into this application in order to more fully describe the state of the art to which this invention pertains.

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Although the present process has been described with reference to specific details of certain embodiments thereof, it is not intended that such details should be regarded as limitations upon the scope of the invention except as and to the extent that they are included in the accompanying claims.

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#### What is claimed is:

- 1. A nucleic acid vector comprising a pair of adeno-associated virus 5 (AAV5) inverted terminal repeats and a promoter between the inverted terminal repeats.
- 2. The vector of claim 1, wherein the promoter is an AAV promoter p5.
- 3. The vector of claim 1, wherein the p5 promoter is AAV5 p5 promoter.
- 4. The vector of claim 1, further comprising an exogenous nucleic acid functionally linked to the promoter.
- 5. The vector of claim 1 encapsidated in an adeno-associated virus particle.
- 6. The particle of claim 5, wherein the particle is an AAV5 particle.
- 7. The particle of claim 5, wherein the particle is an AAV1 particle, an AAV2 particle, an AAV3 particle, an AAV4 particle or an AAV6 particle.
- 8. A recombinant AAV5 virion containing a vector comprising a pair of AAV5 inverted terminal repeats.
- 9. The virion of claim 8, wherein the vector further comprises an exogenous nucleic acid inserted between the inverted terminal repeats.
- An isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO:1.
- 11. An isolated nucleic acid consisting essentially of the nucleotide sequence set forth in SEQ ID NO:1.

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- 12. An isolated nucleic acid that selectively hybridizes with the nucleic acid of claim 11.
- 13. An isolated nucleic acid encoding an adeno-associated virus 5 Rep protein.
- 14. The nucleic acid of claim 13, wherein the adeno-associated virus 5 Rep protein has the amino acid sequence set forth in SEQ ID NO:2.
- 15. The nucleic acid of claim 13, wherein the adeno-associated virus 5 Rep protein has the amino acid sequence set forth in SEQ ID NO:3.
- 16. The nucleic acid of claim 13, wherein the adeno-associated virus 5 Rep protein has the amino acid sequence set forth in SEQ ID NO:12.
- 17. The nucleic acid of claim 13, wherein the adeno-associated virus 5 Rep protein has the amino acid sequence set forth in SEQ ID NO:14.
- 18. An isolated AAV Rep protein.
- 19. The isolated AAV5 Rep protein of claim 18, having the amino acid sequence set forth in SEQ ID NO:2, or a unique fragment thereof.
- 20. The isolated AAV5 Rep protein of claim 18, having the amino acid sequence set forth in SEQ ID NO:3, or a unique fragment thereof.
- 21. An isolated antibody that specifically binds the protein of claim 18.
- 22. An isolated AAV5 capsid protein.
- 23. The isolated AAV5 capsid protein of claim 22 having the amino acid sequence set forth in SEQ ID NO:4.

- 24. An isolated antibody that specifically binds the protein of claim 23.
- 25. The isolated AAV5 capsid protein of claim 22, having the amino acid sequence set forth in SEQ ID NO:5.
- 26. An isolated antibody that specifically binds the protein of claim 25.
- 27. The isolated AAV5 capsid protein of claim 22, having the amino acid sequence set forth in SEQ ID NO:6.
- 28. An isolated antibody that specifically binds the protein of claim 27.
- 29. An isolated nucleic acid encoding the protein of claim 22.
- 30. The nucleic acid of claim 29, having the nucleic acid sequence set forth in SEQ ID NO:7.
- 31. The nucleic acid of claim 29, having the nucleic acid sequence set forth in SEQ ID NO:8.
- 32. The nucleic acid of claim 29, having the nucleic acid sequence set forth in SEQ ID NO:9.
- 33. An isolated nucleic acid that selectively hybridizes with the nucleic acid of claim 29.
- 34. An AAV5 particle comprising a capsid protein consisting essentially of the amino acid sequence set forth in SEQ ID NO:6.
- 35. An isolated nucleic acid comprising an AAV5 p5 promoter.

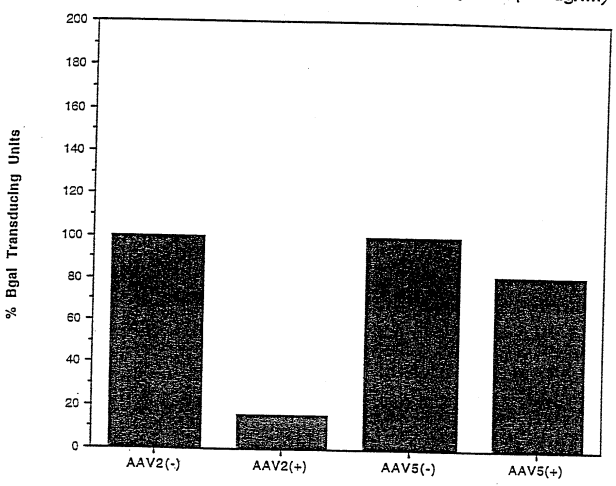
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- 36. A method of screening a cell for infectivity by AAV5, comprising contacting the cell with AAV5 and detecting the presence of AAV5 in the cells.
- 37. A method of determining the suitability of an AAV5 vector for administration to a subject, comprising contacting an antibody-containing sample from the subject with an antigenic fragment of a protein of claim 22 and detecting an antibody-antigen reaction in the sample, the presence of a neutralizing reaction indicating the AAV5 vector to be unsuitable for use in the subject.
- 38. A method of determining the presence in a subject of an AAV5-specific antibody comprising, contacting an antibody-containing sample from the subject with an antigenic fragment of the protein of claim 22 and detecting an antibody-antigen reaction in the sample, the presence of a reaction indicating the presence of an AAV5-specific antibody in the subject.
- 39. A method of delivering a nucleic acid to a cell, comprising administering to the cell an AAV5 particle containing a vector comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to the cell.
- 40. The method of claim 39, wherein the AAV inverted terminal repeats are AAV5 inverted terminal repeats.
- 41. A method of delivering a nucleic acid to a subject comprising administering to a cell from the subject an AAV5 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, and returning the cell to the subject, thereby delivering the nucleic acid to the subject.
- 42. A method of delivering a nucleic acid to a cell in a subject comprising administering to the subject an AAV5 particle comprising the nucleic acid inserted between a pair of AAV inverted terminal repeats, thereby delivering the nucleic acid to a cell in the subject.

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- 43. A method of delivering a nucleic acid to a cell in a subject having antibodies to AAV2 comprising administering to the subject an AAV5 particle comprising the nucleic acid, thereby delivering the nucleic acid to a cell in the subject.
- 44. An isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO:21.
- 45. An isolated nucleic acid comprising the nucleotide sequence set forth in SEQ ID NO: 23.

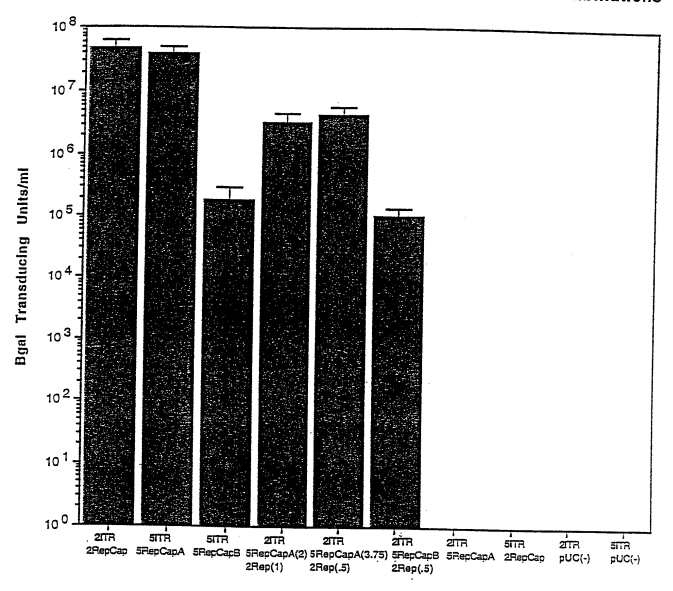
AAV Types 2 & 5 % Inhibition + Heparin (20 ug/ml)



AAV2 or 5, + and - Heparin

FIG. 1

AAV Types 2 & 5 Vector and Helper Plasmid Combinations



AAV2 & 5 Plasmid Combinations

FIG. 2

AAV Types 2 & 5 Tissue Tropism

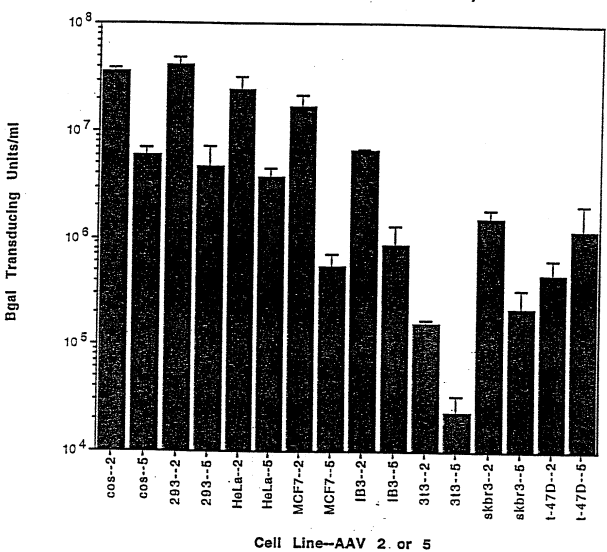


FIG. 3

Apical transduction of human airway epithelia with rAAV2 and rAAV5

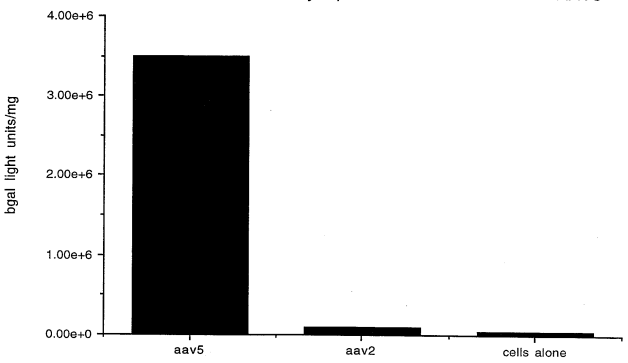


FIG. 7

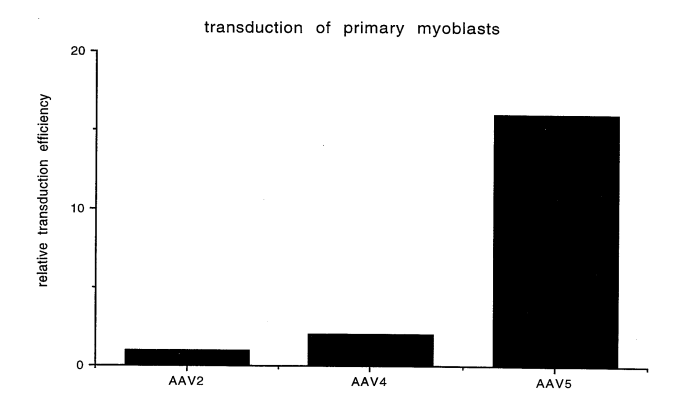


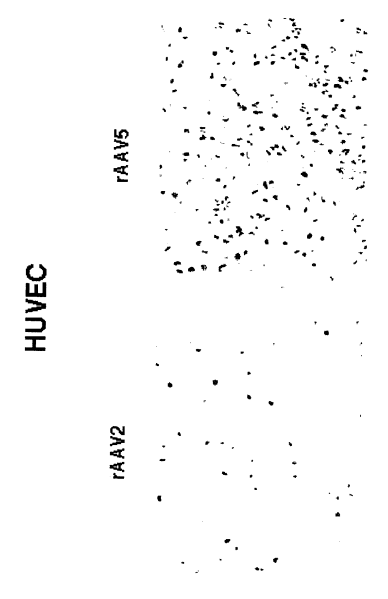
FIG. 8

rAAV5 Primary Rat Brain Explant



FIG. 9





====26-MAY-1999==================================	1E
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The two sequences to be aligned are:	
AAV2CG. Total number of bases: 4679.	
AAV5CG. Total number of bases: 4652.	
Open gap cost : 10 Unit gap cost : 12	
The character to show that two aligned residues are identical is ':'	
AAV2CG - TTGGCCACTCCCTCTCTGGGCGCCTCGCTCGCTCACTGAGGCCGGGCGA -48	
AAV5CG - TGGCACTCTCCCCCTGTCGCGTTCGCTCGCTCGCTCGCTTGGGGGG	
AAV2CG - CCAAAGGTC-GCCCGACGCCCGGGCTTTGCCCGG-GCGGCCTCA	
AAV5CG - CAGCTCAAAGAGCTGCCAGACGACGCCCTCTGGCCGTCGCCCCAAACGAGC -110	
AAV2CGGTGAGCGAGCGCG-CAGAGAGG-GAGTGGCCAACTCCATCACTAGGGGT -141	
AAV5CG - CAGCGAGCGAACGCGACAGGGGGGAGAGTGCCACACTCTCAAGCAAG	
AAV2CG - TCCTGGAGGG-GTGGAGTCGTGACG-TGAATTACGTCATAGGGTTAGGGAGGTCC -194 : :: : : : : : : : : : : : : : : : : :	
AAV5CG - TTTTGTAAGCAGTGATGTCATAATGATGTTAATGCTTATTGTCACGCGATAGTTAA -220	
AAV2CG - TGTATTAGAGGTCACGTGA-GTGTTTTGCGACATTTTGCGACACCATGT -242	
AAV5CG - TG-ATTAACAGTCATGTGATGTGTTTTATCCAATAGGAAGAAGCGCGCGTATGA -274	
AAV2CG - GGTCACGCTGGGTATTTAAGCCCGAGTGAGCACGCAGGGTCTCCAT -288	
AAV5CG - GTTCTCGCGAGACTTCCGGGGTATAAAAGACCGAGTGAACGAGCCCGC-CGCCAT -328	
AAV2CG - T-TTGAAGCGGGAG-GTTTGAACGCGCA-GCCGCCATGCCGGGGTTTTACGAGAT -340	
AAV5CG - TCTTTGCTCTGGACTGCTAGAGGACCCTCGCTGCCATGGCTACCTTCTATGAAGT -383	
AAV2CG - TGTGATTAAGGTCCCCAGCGACCTTGACGGGCATCTGCCCGGCATTTCTGACAGC -395	
: :: ::::: :::::::::::::::::::::::::::	
AAV2CG - TTTGTGAACTGGGTGGCCGAGAAGGAATGGGAGTTGCCGCCAGATTCTGACATGG -450	
AAV5CG - TTTGTGGACTGGGTAACTGGTCAAATTTGGGAGCTGCCTCCAGAGTCAGATTTAA -493	

FIG. 4

AAV2CG	- ATCTGAATCTC L'TGAGCAGGCACCCCTGACCGTGGCCGAGACTGCAGCGCGA -505
AAV5CG	- ATTTGACTCTGGTTGAACAGCCTCAGTTGACGGTGGCTGATAGAATTCGCCGCGT -548
AAV2CG	- CTTTCTGACGGAATGGCGCCGTGTGAGTAAGGCCCCGGAGGCCCTTTTCTTTGTG -560
AAV5CG	- GTTCCTGTACGAGTGGAACAAATTTTCCAAGCAGGAGTCCAAATTCTTTGTG -600
AAV2CG	- CAATTTGAGAAGGGAGAGAGCTACTTCCACATGCACGTGCTCGTGGAAACCACCG -615
AAV5CG	- CAGTTTGAAAAGGGATCTGAATATTTTCATCTGCACACGCTTGTGGAGACCTCCG -655
AAV2CG	- GGGTGAAATCCATGGTTTTGGGACGTTTCCTGAGTCAGATTCGCGAAAAACTGAT -670
AAV5CG	- GCATCTCTTCCATGGTCCTCGGCCGCTACGTGAGTCAGATTCGCGCCCAGCTGGT -710
AAV2CG	- TCAGAGAATTTACCGCGGGATCGAGCCGACTTTGCCAAACTGGTTCGCGGTCACA -725
AAV5CG	- GAAAGTGGTCTTCCAGGGAATTGAACCCCAGATCAACGACTGGGTCGCCATCACC -765
AAV2CG	- AAGACCAGAAATGGCGCCGGAGGCGGGAACAAGGTGGTGGATGAGTGCTACATCC -780
AAV5CG	- AAGGTAAAGAAGGGCGGAGCCAATAAGGTGGTGGATTCTGGGTATATTC -814
AAV2CG	- CCAATTACTTGCTCCCCAAAACCCAGCCTGAGCTCCAGTGGGCGTGGACTAATAT -835
AAV5CG	- CCGCCTACCTGCTGCCGAAGGTCCAACCGGAGCTTCAGTGGGCGTGGACAAACCT -869
AAV2CG	- GGAACAGTATTTAAGCGCCTGTTTGAATCTCACGGAGCGTAAACGGTTGGTGGCG -890
AAV5CG	- GGACGAGTATAAATTGGCCGCCCTGAATCTGGAGGAGCGCAAACGGCTCGTCGCG -924
AAV2CG	- CAGCATCTGACGCACGTGTCGCAGACGAGCAGGAGCAGAACAAAGAGAATCAGAATC -945
AAV5CG	- CAGTTTCTGGCAGAATCCTCGCAG-CGCTCGCAGGAGGCGGCTTCGCAGCGTG -976
AAV2CG	- CCAATTCTGATGCGCCGGTGATCAGATCAAAAACTTCAGCCAGGTACATGGAGCT -1000
AAV5CG	- AGTTCTCGGCTGACCCGGTCATCAAAAGCAAGACTTCCCAGAAATACATGGCGCT -1031
AAV2CG	- GGTCGGGTGGCTCGTGGACAAGGGGATTACCTCGGAGAAGCAGTGGATCCAGGAG -1055
AAV5CG	- CGTCAACTGGCTCGTGGAGCACGGCATCACTTCCGAGAAGCAGTGGATCCAGGAA -1086
AAV2CG	- GACCAGGCCTCATACATCTCCTTCAATGCGGCCTCCAACTCGCGGTCCCAAATCA -1110
AAV5CG	- AATCAGGAGAGCTACCTCCTTCAACTCCACCGGCAACTCTCGGAGCCAGATCA -1141
AAV2CG	- AGGCTGCCTTGGACAATGCGGGAAAGATTATGAGCCTGACTAAAACCGCCCCCGA -1165
AAV5CG	- AGGCCGCGCTCGACAACGCGACCAAAATTATGAGTCTGACAAAAAGCGCGGTGGA -1196
AAV2CG	- CTACCTGGTGGGCCAGCAGCCCGTG-GAGGACATTTCCAGCAATCGGATTTATAA -1219
AAV5CG	- CTACCTCGTGGGG-AGCTCCGTTCCCGAGGACATTTCAAAAAACAGAATCTGGCA -1250

AAV2CG	- AATTTTGGAACTAAACGGGTACGATCCCCAATATGCGGCTTCCGTCTTTCTGGGA -1274
AAV5CG	- AATTTTTGAGATGAATGGCTACGACCCGGCCTACGCGGGATCCATCC
AAV2CG	- TGGGCCACGAAAAAGTTCGGCAAGAGGAACACCATCTGGCTGTTTGGGCCTGCAA -1329
AAV5CG	- TGGTGTCAGCGCTCCTTCAACAAGAGGAACACCGTCTGGCTCTACGGACCCGCCA -1360
AAV2CG	- CTACCGGGAAGACCAACATCGCGGAGGCCATAGCCCACACTGTGCCCTTCTACGG -1384
AAV5CG	- CGACCGGCAAGACCAACATCGCGGAGGCCATCGCCCACACTGTGCCCTTTTACGG -1415
AAV2CG	- GTGCGTAAACTGGACCAATGAGAACTTTCCCTTCAACGACTGTGTCGACAAGATG -1439
AAV5CG	- CTGCGTGAACTGGACCAATGAAAACTTTCCCTTTAATGACTGTGTGGACAAAATG -1470
AAV2CG	- GTGATCTGGTGGGAGGGGGAAGATGACCGCCAAGGTCGTGGAGTCGGCCAAAG -1494
AAV5CG	: :: :::::::::::::::::::::::::::::::::
AAV2CG	- CCATTCTCGGAGGAAGCAAGGTGCGCGTGGACCAGAAATGCAAGTCCTCGGCCCA -1549
AAV5ÇG	- CCATCCTGGGGGGCTCAAAGGTGCGGGTCGATCAGAAATGTAAATCCTCTGTTCA -1580
AAV2CG	- GATAGACCCGACTCCCGTGATCGTCACCTCCAACACCAACATGTGCGCCGTGATT -1604
AAV5CG	::::::::::::::::::::::::::::::::::::::
AAV2CG	- GACGGGAACTCAACGACCTTCGAACACCAGCAGCCGTTGCAAGACCGGATGTTCA -1659
AAV5CG	- GATGGGAATTCCACGACCTTTGAACACCAGCAGCCGCTGGAGGACCGCATGTTCA -1690
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AAV2CG	- AGTCAAAGACTTTTTCCGGTGGGCAAAGGATCACGTGGTTGAGGTGGAGCATGAA -1769
AAV5CG	- AGTCAAGGACTTTTTGCTTGGGCAAAGGTCAATCAGGTGCCGGTGACTCACGAG -1800
AAV2CG	- TTCTACGTCAAAAAGGGTGGAGCCAAGAAAAGACCCGCCCCAGTGACGCAGA -1822
AAV5CG	:: ::: ::: ::: ::: ::: ::: ::: - TTTAAAGTTCCCAGGGAATTGGCGGGAACTAAAGGGGCGGAGAAATCTC -1849
AAV2CG	- TATAAGTGAGCCCAAACGGGTGCGCGAGTCAGTTGCGCAGCCATCGACGTCAGAC -1877
AAV5CG	:: :: ::::::::::::::::::::::::::::::::
AAV2CG	- GCGGAAGCTTCGATCAACTACGCAGACAGGTACCAAAACAAAT-GTTCTCGTCAC -1931
AAV5CG	: :::: : :: :: :: :: :: :: :: :: :: ::
AAV2CG	- GTGGGCATGAATCT-GATGCTGTTTCCCTGCAGACAATGCGAGAGAATGAATCAG -1985

AAV5CG	- GTGACTGTTGA JCCGCTCCTCTGCGACCGCTCA-ATTGGAA.1CAAGGTATG -1999
AAV2CG	- AATTCAAATATCTGCTTCACTCACGGACAGAAAGACTGTTTAGAGTGCTTTCCCG -2040
AAV5CG	: ::::: :: :: :: :: :: :: :: :: :: :: :
AAV2CG	- TGTCA-GAATCTCAACCCGTTTCTGTCGTCAAAAAGGCGTATCAGAAACTGTG -2092
AAV5CG	- TGTGATGAATGTGAATATTTGAATCGGGGCAAAAATGGATGTATCTGTCACAATG -2101
AAV2CG	- CTACATTCA-TCATATCATGGGAAAGGTGCCAGACGCTTGCACTGCCTGCG -2142
AAV5CG	- TAACTCACTGTCAAATTTGTCATGGGATTCCCCCCTGGGAAAAGGAAAACTTG2154
AAV2CG	- ATCTGGTCAATGTGGATTTGGATGACTGCATCTTTGAACAATAAATGATTTAAAT -2197
AAV5CG	TCAGATTT-TGGGGATTTTGACGATGCCAATAAAGAACAGTAAATAAAGCGAGT -2207
AAV2CG	- CAGGTATGGCTGCCGATGGTTATCTTCCAGATTGGCTCGAGGACACTCTCTCT
AAV5CG	AGTCATGTCTTTGTTGATCACCCTCCAGATTGGTTGGAAGAAGTTGGTGA -2258
AAV2CG	- AGGAATAAGACAGTGGTGGAAGCTCAAACCTGGCCCACCACCACCACCAAAGCCCGCA -2307
AAV5CG	- AGGTCTTCGCGAGTTTTTGGGCCTTGAAGCGGGCCCACCGAAACCAAAACCCAAT -2313
AAV2CG	- GAGCGGCATAAGGACGACAGCAGGGGTCTTGTGCTTCCTGGGTACAAGTACCTCG -2362
AAV5CG	- CAGCAGCATCAAGATCAAGCCCGTGGTCTTGTGCTGCCTGGTTATAACTATCTCG -2368
AAV2CG	- GACCCTTCAACGGACTCGACAAGGGAGAGCCGGTCAACGAGGCAGACGCCGCGGC -2417
AAV5CG	- GACCCGGAAACGGTCTCGATCGAGGAGAGCCTGTCAACAGGGCAGACGAGGTCGC -2423
AAV2CG	- CCTCGAGCACAAAGCCTACGACCGGCAGCTCGACAGCGGAGACAACCCGTAC -2472
AAV5CG	- GCGAGAGCACGTCCGTACAACGAGCAGCTTGAGGCGGGAGACAACCCCTAC -2478
AAV2CG	- CTCAAGTACAACCACGCCGACGCGGAGTTTCAGGAGCGCCTTAAAGAAGATACGT -2527
AAV5CG	- CTCAAGTACAACCACGCGGACGCCGAGTTTCAGGAGAAGCTCGCCGACGACACAT -2533
AAV2CG	- CTTTTGC GGCAACCTCGGACGAGCAGTCTTCCAGGCGAAAAAGAGGGGTTCTTGA -2582
AAV5CG	- CCTTCGGGGGAAACCTCGGAAAGGCAGTCTTTCAGGCCAAGAAAAGGGTTCTCGA -2588
AAV2CG	- ACCTCTGGGCCTGGTTGAGGAACCTGTTAAGACGGCTCCGGGAAAAAAGAGGCCG -2637
AAV5CG	- ACCTTTTGGCCTGGTTGAAGAGGGTGCTAAGACGGCCCCTACCGGAAAGCGGATA -2643
AAV2CG	- GTAGAGCACTCTCCTGTGGAGCCAGACTCCTCCTCGGGAACCGGAAAGGCGGGCC -2692
AAV5CG	: :: ::: ::: ::: ::: ::: ::: ::: ::: :
AAV2CG	- AGCAGCCTGCAAGAAAAAGATTGAATTTTGGTCAGACTGGAGACGCAG-ACTCAG -2746

AAV5CG	: :::::: : :: :::::: : :::::::::::::::
AAV2CG	- TACCTGACCCCCAGCCTCTCGGACAGCCACCAGCAGCCCCCTCTGGTCTGGGAAC -2801
AAV5CG	CGGATCCC-AGCAGCTGCAAATCCCAGCCCAACCAGCCTCAAGTTTGGGAGC -2780
AAV2CG	- TAATACGATGGCTACAGGCAGTGGCGCACCAATGGCAGACAATAACGAGGGCGCC -2856
AAV5CG	- TGATACAATGTCTGCGGGAGGTGGCGGCCCATTGGGCGACAATAACCAAGGTGCC -2835
AAV2CG	- GACGGAGTGGGTAATTCCTCGGGAAATTGGCATTGCGATTCCACATGGATGG
AAV5CG	- GATGGAGTGGGCAATGCCTCGGGAGATTGGCATTGCGATTCCACGTGGATGGGGG -2890
AAV2CG	- ACAGAGTCATCACCACCAGCACCCGAACCTGGGCCCTGCCCACCTACAACAACCA -2966
AAV5CG	- ACAGAGTCGTCACCAAGTCCACCCGAACCTGGGTGCTGCCCAGCTACAACAACCA -2945
AAV2CG	- CCTCTACAAACAAATTTCCAGCCAATCAGGAGGGTCGAACGACAATCACTAC =3018
AAV5CG	- CCAGTACCGAGAGATCAAAAGCGGCTCCGTCGACGGAAGCAACGCCAACGCCTAC -3000
AAV2CG	- TTTGGCTACAGCACCCCTTGGGGGTATTTTGACTTCAACAGATTCCACTGCCACT -3073
AAV5CG	- TTTGGATACAGCACCCCTGGGGGTACTTTGACTTTAACCGCTTCCACAGCCACT -3055
AAV2CG	- TTTCACCACGTGACTGGCAAAGACTCATCAACAACAACTGGGGATTCCGACCCAA -3128
AAV5CG	- GGAGCCCCGAGACTGGCAAAGACTCATCAACAACTACTGGGGCTTCAGACCCCG -3110
AAV2CG	- GAGACTCAACTTCAAGCTCTTTAACATTCAAGTCAAAGAGGTCACGCAGAATGAC -3183
AAV5CG	- GTCCCTCAGAGTCAAAATCTTCAACATTCAAGTCAAAGAGGTCACGGTGCAGGAC -3165
AAV2CG	- GGTACGACGACTGCCAATAACCTTACCAGCACGGTTCAGGTGTTTACTGACT -3238
AAV5CG	- TCCACCACCACCACCACCACCACCTCACCTCCACCGTCCAAGTGTTTACGGACG -3220
AAV2CG	- CGGAGTACCAGCTCCCGTACGTCCTCGGCTCGGCGCATCAAGGATGCCTCCCGCC -3293
AAV5CG	- ACGACTACCAGCTGCCCTACGTCGGCAACGGGACCGAGGGATGCCTGCC
AAV2CG	- GTTCCCAGCAGACGTCTTCATGGTGCCACAGTATGGATACCTCACCCTGAACAAC -3348
AAV5CG	- CTTCCCTCCGCAGGTCTTTACGCTGCCGCAGTACGGTTACGCGACGCTGAACCGC -3330
AAV2CG	- GGGAGT-CAGGCAGTAGGACGCTCTTCATTTTACTGCCTGGAGTACTTTC -3397
AAV5CG	: ::::::::::::::::::::::::::::::::::::
AAV2CG	- CTTCTCAGATGCTGCGTACCGGAAACAACTTTACCTTCAGCTACACTTTTGAGGA -3452
AAV5CG	: ::::::::::::::::::::::::::::::::::::

AAV2CG	- CGTTCCTTTCL_JAGCAGCTACGCTCACAGCCAGAGTCTGGALJGTCTCATGAAT -3507
AAV5CG	- GGTGCCCTTCCACTCCAGCTTCGCTCCCAGTCAGAACCTGTTCAAGCTGGCCAAC -3495
AAV2CG	- CCTCTCATCGACCAGTACCTGTATTACTTGAGCAGAACAACACTC3553
AAV5CG	- CCGCTGGTGGACCAGTACTTGTACCGCTTCGTGAGCACAAATAACACTGGCGGAG -3550
AAV2CG	CAAGTGGAACCACCACGCAGTCA-AGGCTTCAGTTTTCTCAGGCCGGAG -3601
AAV5CG	- TCCAGTTCAACAAGAACCTGGCCGGGAGATACGCCAACACCTACAAAAACTGGTT -3605
AAV2CG	- CGAGTGACATTCGGGACCAGTCTAGGAACTGGCTTCCTGGACCCTGTTACCGCCA -3656
AAV5CG	- CCCGGGGCCCATGGGCCCGAACCCAGGG-CTGGAA-CCTGGGCTCCGGGGTCAACC -3658
AAV2CG	- GCAGCGAGTATCAAAGACATCTGCGGATAACAACAACAGTGAATACTCGTGGACT -3711
AAV5CG	- GC-GCCAGTGTCAGCGCCTTC-GCCACGACCAATAGGA-TGGAG-CTCGAGGGCG -3709
AAV2CG	- GGAGCTACCAAGTACCACCTCAATGGCAGAGACTCTCTGGTGAATCCGGGCCCGG -3766
AAV5CG	- CGAGTTACCAGGTGCCCCCGCAGCCGA-ACGGCATGACCAACAACCTCCAGG -3760
AAV2CG	- CCATGGCAAGCCACAAGGACGATGAAGAAAGTTTTTTCCTCAGAGCGGGGTTCT -3821
AAV5CG	- GCAGCAACACCTATGCCCTGGAGAACACTATGATCTTCAACAGC3804
AAV2CG	- CATCTTTGGGAAGCAAGGCTCAGAGAAAACAAATGTGGACATTGAAAAGGTCATG -3876
AAV5CG	- CAGCCG-GCGAACCCGGGCACCACGCCACGTACCTCGAGGGCAACATGCTCATC -3858
AAV2CG	- ATTACAGACGAAGAGAAATCAGGACAACCAATCCCGTGGC-TACGGAGCAGTAT -3930
AAV5CG	- ACCAG-CGAGAGCGAGACGCAGCCGGTGAACCGCGTGCGTACAACGTCGGCG -3910
AAV2CG	- GGTTCTGTATCTACCAACCTCCAGAGAGGCAACAGACAGCAGCTACCGCAGATG -3985
AAV5CG	- GGCAGA-TGGCCACCAACAACCAGAGCTCCACCACTGCCCCCGCGACCGGCACGT -3964
AAV2CG	- TCAACACACAAGGCGTTCTTCCAGGCATGGTCTGGCAGGACAGAGATGTGTACCT -4040
AAV5CG	- ACAACCTCCAGGAAATCGTGCCCGGCAGCGTGTGGATGGA
AAV2CG	- TCAGGGGCCCATCTGGGCAAAGATTCCACACACGGACGACATTTTCACCCCTCT -4095
AAV5CG	- CCAAGGACCCATCTGGGCCAAGATCCCAGAGACGGGGGGGCGCACTTTCACCCCTCT -4074
AAV2CG	- CCCCTCATGGGTGGATTCGGACTTAAACACCCTCCTCCACAGATTCTCATCAAGA -4150
AAV5CG	- CCGGCCATGGGCGGATTCGGACTCAAACACCCACCGCCCATGATGCTCATCAAGA -4129
AAV2CG	- ACACCCGGTACCTGCGAATCCTTCGACCACCTTCAGTG-CGGCAAAGTTTGCTT -4204
AAV5CG	- ACACGCCTGTGCCCGGAAATATC-ACCAGCTTCTCGGACGTGCCCGTCAGCAG -4181

```
- CCTTCATCACACAGTACTCCACGGGACAGGTCAGCGTGGAGATCGAGTGGGAGCT -4259
 AAV2CG
         - C-TTCATCACCCAGTACAGCACCGGGCAGGTCACCGTGGAGATGGAGTGGGAGCT -4235
 AAV5CG
        - GCAGAAGGAAAACAGCAAACGCTGGAATCCCGAAATTCAGTACACTTCCAACTAC -4314
 AAV2CG
          - CAAGAAGGAAAACTCCAAGAGGTGGAACCCAGAGATCCAGTACACAACAACTAC -4290
 AAV5CG
       - AACAAGTCTGTTAATGTGGACTTTACTGTGGACACTAATGGCGTGTATTCAGAGC -4369
AAV2CG
         - AACGACCCCAGTTTGTGGACTTTGCCCCGGACAGCACCGGGGA--ATACAGAAC -4343
AAV5CG
       - CTC--GCCCCATTGGCACCAGATACCTGACTCGTAATCTGTAAT---TGCTTGT- -4418
AAV2CG
         - CACCAGACCTATCGGAACCCGATACCTTACCCGACCCCTTTAACCCATTCATGTC -4398
AAV5CG
       - ---TAA---TCAATAAACCGTTTAATTCGTTTCAGTTGAACTTTGG-TCTCTGCGT -4467
AAV2CG
          - GCATACCCTCAATAAACCGTGTA-TTCGTGTCAGTAAAATACTGGGTCTTGTGGT -4452
AAV5CG
       - ATTTCTTTCT-TATCTAGTTTCCATGGCTACGTAGATAAGTAGCATGGCGGGTTA -4521
AAV2CG
         AAV5CG
       - ATCATTAACTACAAGGAACCCCTAGTGATGGAGTTGGCCACTCCCTC-TCTGCGC -4575
AAV2CG
         AAV5CG
       - GCTCGCTCGCTCACTGAG--GCCGGGCGACCAAAGGTCGCCCGACGCCCGGGCTT -4628
AAV2CG
        : : :::: :::: :::: :::::
       - GGGTGGCAGCTCAAAGAGCTGCCAGACGACGCCCTCTGGCCGTCGCCCC---- -4604
AAV5CG
       - TGCCCGGGCGCCTCAGTGAGCGAGCGCGCGCAGAGAGGGAGTGGCCAA -4679
AAV2CG
         - --CCCAAACGAGC-CAGCGAGCGAGCGACGCGACAGGGGGGAGAGTGCCA -4652
AAV5CG
Identity : 3013 (64.77%)
Number of gaps inserted in AAV2CG: 43
Number of gaps inserted in AAV5CG: 63
```

<sup>===26-</sup>MAY-1999========PC/GENE===

## FIG. 5

```
=PC/GENE===
                  ----PALIGN=
AN-1997=
**********
MENT OF TWO PROTEIN SEQUENCES. *
*********
                                        THILL ACG
sequences to be aligned are:
1
.V2
umber of residues: 735.
.V5VP1
V5VP1
umber of residues: 724. 2/01-4272
son matrix : Structure-genetic matrix.
          : 8
p cost
          : 5
p cost
racter to show that two aligned residues are identical is ':'
racter to show that two aligned residues are similar is '.'
cids said to be 'similar' are: A,S,T; D,E; N,Q; R,K; I,L,M,V; F,Y,W

    MAADGYLPDWLEDTLSEGIRQWWKLKPGPPPPKPAERHKDDSRGLVLPGYKYLGP -55

         - MSFVDHPPDWLEE-VGEGLREFLGLEAGPPKPKPNQQHQDQARGLVLPGYNYLGP -54
   - FNGLDKGEPVNEADAAALEHDKAYDRQLDSGDNPYLKYNHADAEFQERLKEDTSF -110
     - GNGLDRGEPVNRADEVAREHDISYNEQLEAGDNPYLKYNHADAEFQEKLADDTSF -109
   - GGNLGRAVFQAKKRVLEPLGLVEEPVKTAPGKKRPVEHSPVEPDSSSGTGKAGQQ -165
    - GGNLGKAVFQAKKRVLEPFGLVEEGAKTAPTGKRIDDHFPKR--KKARTEEDSKP -162
   - PARKRLNFGQTGDADSVPDPQPLGQPPAAPSGLGTNTMATGSGAPMADNNEGADG -220
Ţ
                    -----SDAEAGPSGSQQLQIPAQPASSLGADTMSAGGGGPLGDNNQGADG -210
1
   - VGNSSGNWHCDSTWMGDRVITTSTRTWALPTYNNHLYKQISSQSG-ASNDNHYFG -274
1
     - VGNASGDWHCDSTWMGDRVVTKSTRTWVLPSYNNHQYREIKSGSVDGSNANAYFG -265
1
   - YSTPWGYFDFNRFHCHFSPRDWQRLINNNWGFRPKRLNFKLFNIQVKEVTQNDGT -329
1
     - YSTPWGYFDFNRFHSHWSPRDWQRLINNYWGFRPRSLRVKIFNIQVKEVTVQDST -320
   - TTIANNLTSTVQVFTDSEYQLPYVLGSAHQGCLPPFPADVFMVPQYGYLTLNNGS -384
                            :::: :: :: :::::: :::
     - TTIANNLTSTVQVFTDDDYQLPYVVGNGTEGCLPAFPPQVFTLPQYGYATINRDN -375
1
```

- QAVGRSSFYCLEYFPSQMLRTGNNFTFSYTFEDVPFHSSYAHSQSLDRLMNPL	-43/
- TENPTERSSFFCLEYFPSKMLRTGNNFEFTYNFEEVPFHSSFAPSQNLFKLANPL	-430
- IDQYLYYLSRTNTPSGTTTQSRLQFSQAGASDIRDQSRNWLPGPCYRQQRVSKTS	-492
- VDQYLYRFVSTNNTGGVQFNKNLAGRYANTYKNWFPGPMGRTQGWNLGS	-479
- ADNNNSEYSWTGATKYHLNGRDSLVNPGPAMASHKDDEEKFFPQSGVLIFGKQGS	-547
- GVNRASVSAFATTNRMELEGASYQVPPQPNGMTNNLQGSNTYALENTMIFNSQPA	-534
- EKTNVDIEKVMITDEEEIRTTNPVATEQYGSVSTNLQRGNRQAATADVNTQG	-599
- NPGTTATYLEGNMLITSESETQPVNRVAYNVGGQMATNNQSSTTAPATGTYNLQE	<del>-</del> 589
- VLPGMVWQDRDVYLQGPIWAKIPHTDGHFHPSPLMGGFGLKHPPPQILIKNTPVP	-654
- IVPGSVWMERDVYLQGPIWAKIPETGAHFHPSPAMGGFGLKHPPPMMLIKNTPVP	-644
- ANPSTTFSAAKFASFITQYSTGQVSVEIEWELQKENSKRWNPEIQYTSNYNKSVN	-709
- GNI-TSFSDVPVSSFITQYSTGQVTVEMEWELKKENSKRWNPEIQYTNNYNDPQF	-698
- VDFTVDTNGVYSEPRPIGTRYLTRNL -735	
:::. :: ::::::::::::::::::::::::::::::	
. 421 (EQ 159)	
: 421 (58.15%) y: 63 (8.70%)	
gaps inserted in AAV2VP1: 3 gaps inserted in AAV5VP1: 5	
	DC/GENE

# FIG. 6

```
PC/GENE=
7-31-DEC-1996=
*********
* ALIGNMENT OF TWO PROTEIN SEQUENCES. *
***********
The two sequences to be aligned are:
REP78.
   REP78
DE
   AAV
os
Total number of residues: 621.
AAVSREP.
DE
   REP
    AAV5
OS
Total number of residues: 610.
Comparison matrix : Structure-genetic matrix.
               : 8
Open gap cost
               : 5
Unit gap cost
The character to show that two aligned residues are identical is ':'
The character to show that two aligned residues are similar is '.'
Amino acids said to be 'similar' are: A,S,T; D,E; N,Q; R,K; I,L,M,V; F,Y,W
        - MPGFYEIVIKVPSDLDGHLPGISDSFVNWVAEKEWELPPDSDMDLNLIEQAPLTV -55
REP78
          - MATFYEVIVRVPFDVEEHLPGISDSFVDWVTGQIWELPPESDLNLTLVEQPQLTV -55
AAV5REP
        - AEKLORDFLTEWRRVSKAPEALFFVQFEKGESYFHMHVLVETTGVKSMVLGRFLS -110
REP78
          - ADRIRRVFLYEWNKFSKQ-ESKFFVQFEKGSEYFHLHTLVETSGISSMVLGRYVS -109
AAVSREP
        - QIREKLIQRIYRGIEPTLPNWFAVTKTRNGAGGGNKVVDECYIPNYLLPKTQPEL -165
        - QIRAQLVKVVFQGIEPQINDWVAITKVKKG--GANKVVDSGYIPAYLLPKVQPEL -162
REP78
AAV5REP
        - QWAWTNMEQYLSACLNLTERKRLVAQHLTHVSQTQEQNKENQNPNSDAPVIRSKT -220
REP78
          - QWAWTNLDEYKLAALNLEERKRLVAQFLA-ESSQRSQEAASQREFSADPVIKSKT -216
AAV5REP
        - SARYMELVGWLVDKGITSEKQWIQEDQASYISFNAASNSRSQIKAALDNAGKIMS -275
REP78
          - SQKYMALVNWLVEHGITSEKQWIQENQESYLSFNSTGNSRSQIKAALDNATKIMS -271
AAV5REP
        - LTKTAPDYLVGQQPVEDISSNRIYKILELNGYDPQYAASVFLGWATKKFGKRNTI -330
REP78
                      - LTKSAVDYLVGSSVPEDISKNRIWQIFEMNGYDPAYAGSILYGWCQRSFNKRNTV -326
 AAVSREP
         - WLFGPATTGKTNIAEAIAHTVPFYGCVNWTNENFPFNDCVDKMVIWWEEGKMTAK -385
 REP78
          - WLYGPATTGKTNIAFAIAHTVPFYGCVNWTNENFPFNDCVDKMLIWWEEGKMTNK -381
 AAV5REP
```

===31-DEC-1996==

PC/GENE=

```
- VVESAKAILGGSKVRVDQKCKSSAQIDPTPVIVTSNTNMCAVIDGNSTTFEHQQP -440
REP78
          - VVESAKAILGGSKVRVDQKCKSSVQIDSTPVIVTSNTNMCVVVDGNSTTFEHQQP -436
AAV5REP
          - LQDRMFKFELTRRLDHDFGKVTKQEVKDFFRWAKDHVVEVEHEFYVKKGGAKKRP -495
REP78
            - LEDRMFKFELTKRLPPDFGKITKQEVKDFFAWAKVNQVPVTHEFKVPRELAGTK- -490
AAV5REP
          - APSDADISEPKRVRESVAQPSTSDAFASINYADRYQNKCSRHVGMNLMLFFCRQC -550
REP78
                       : .
                                   : :
          - GAEKSLKRPLGDVTNTXYKSLEKRARLSFVPETPRSSDVTVDPAPLRPLNWNSRY -545
AAV5REP
          - ERMNQNSNICFTHGQKDCLECFPVSESQPVSVVKKAYQKLCYIHHIMGKVPDACT -605
REP78
                            : ::
           DC-KCDYHAQFDNISNKCDECEYINRGKNGCICHNVTH-CQICHGIPPWEKENLS -598
AAV5REP
          - ACDLVNVDLDDCIFEQ -621
REP78
                  ::: ::
AAV5REP
          - DF----GDFDDANKEQ -610
Identity : 355 ( 58.2%)
Similarity: 56 ( 9.2%)
Number of gaps inserted in REP78: 0
Number of gaps inserted in AAV5REP: 7
```

1

## SEQUENCE LISTING

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<110> Chiorini, John
      <120> AAV5 VECTOR AND USES THEREOF
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      <150> 60/087,029
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      <211> 4652
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                                                                       120
gcgaacgcga caggggggag agtgccacac tctcaagcaa gggggttttg taagcagtga
                                                                       180
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                                                                       240
tgttttatcc aataggaaga aagcgcgcgt atgagttctc gcgagacttc cggggtataa
                                                                       300
aagaccgagt gaacgagccc gccgccattc tttgctctgg actgctagag gaccctcgct
                                                                       360
gccatggcta ccttctatga agtcattgtt cgcgtcccat ttgacgtgga ggaacatctg
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gagtcagatt taaatttgac tctggttgaa cagcctcagt tgacggtggc tgatagaatt
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                                                                       600
cagtttgaaa agggatctga atattttcat ctgcacacgc ttgtggagac ctccggcatc
                                                                       660
tettecatgg teeteggeeg etacgtgagt cagattegeg eccagetggt gaaagtggte
                                                                       720
ttccagggaa ttgaacccca gatcaacgac tgggtcgcca tcaccaaggt aaagaagggc
                                                                       780
ggagccaata aggtggtgga ttctgggtat attcccgcct acctgctgcc gaaggtccaa
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                                                                     1680
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PCT/US99/11958

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ttcaagctgg	ccaacccgct	ggtggaccag	tacttgtacc	gcttcgtgag	cacaaataac	3540
actggcggag	tccagttcaa	caagaacctg	gccgggagat	acgccaacac	ctacaaaaac	3600
tggttcccgg	ggcccatggg	ccgaacccag	ggctggaacc	tgggctccgg	ggtcaaccgc	3660
gccagtgtca	gcgccttcgc	cacgaccaat	aggatggagc	tcgagggcgc	gagttaccag	3720
gtgcccccgc	agccgaacgg	catgaccaac	aacctccagg	gcagcaacac	ctatgccctg	3780
gagaacacta	tgatcttcaa	cagccagccg	gcgaacccgg	gcaccaccgc	cacgtacctc	3840
gagggcaaca	tgctcatcac	cagcgagagc	gagacgcagc	cggtgaaccg	cgtggcgtac	3900
aacgtcggcg	ggcagatggc	caccaacaac	cagageteca	ccactgcccc	cgcgaccggc	3960
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caaggaccca	tctgggccaa	gatcccagag	acgggggcgc	actttcaccc	ctctccggcc	4080
atgggcggat	tcggactcaa	acacccaccg	cccatgatgc	tcatcaagaa	cacgcctgtg	4140
cccggaaata	tcaccagctt	ctcggacgtg	cccgtcagca	gcttcatcac	ccagtacagc	4200
accgggcagg	tcaccgtgga	gatggagtgg	gagctcaaga	aggaaaactc	caagaggtgg	4260
aacccagaga	tccagtacac	aaacaactac	aacgaccccc	agtttgtgga	ctttgccccg	4320
gacagcaccg	gggaatacag	aaccaccaga	cctatcggaa	cccgatacct	tacccgaccc	4380
ctttaaccca	ttcatgtcgc	ataccctcaa	taaaccgtgt	attcgtgtca	gtaaaatact	4440
gcctcttgtg	gtcattcaat	gaataacagc	ttacaacatc	tacaaaacct	ccttgcttga	4500
gagtgtggca	ctctccccc	tgtcgcgttc	gctcgctcgc	tggctcgttt	ggggggtgg	4560
cagctcaaag	agctgccaga	cgacggccct	ctggccgtcg	ccccccaaa	cgagccagcg	4620
agcgagcgaa	cgcgacaggg	gggagagtgc	ca			4652

<210> 2

<211> 390

<212> PRT

<213> Artificial Sequence

<220>

<400> 2

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Gln Trp Ile Gln Glu Asn Gln Glu Ser Tyr Leu Ser Phe Asn Ser Thr
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Gly Asn Ser Arg Ser Gln Ile Lys Ala Ala Leu Asp Asn Ala Thr Lys
                          40
Ile Met Ser Leu Thr Lys Ser Ala Val Asp Tyr Leu Val Gly Ser Ser
Val Pro Glu Asp Ile Ser Lys Asn Arg Ile Trp Gln Ile Phe Glu Met
                                      75
Asn Gly Tyr Asp Pro Ala Tyr Ala Gly Ser Ile Leu Tyr Gly Trp Cys
                                  90
Gln Arg Ser Phe Asn Lys Arg Asn Thr Val Trp Leu Tyr Gly Pro Ala
                    105
Thr Thr Gly Lys Thr Asn Ile Ala Glu Ala Ile Ala His Thr Val Pro
       115
                120
Phe Tyr Gly Cys Val Asn Trp Thr Asn Glu Asn Phe Pro Phe Asn Asp
            135
Cys Val Asp Lys Met Leu Ile Trp Trp Glu Glu Gly Lys Met Thr Asn
                  150
                                      155
Lys Val Val Glu Ser Ala Lys Ala Ile Leu Gly Gly Ser Lys Val Arg
                                  170
Val Asp Gln Lys Cys Lys Ser Ser Val Gln Ile Asp Ser Thr Pro Val
                              185
Ile Val Thr Ser Asn Thr Asn Met Cys Val Val Val Asp Gly Asn Ser
                          200
Thr Thr Phe Glu His Gln Gln Pro Leu Glu Asp Arg Met Phe Lys Phe
                       215
Glu Leu Thr Lys Arg Leu Pro Pro Asp Phe Gly Lys Ile Thr Lys Gln
                   230
                                      235
Glu Val Lys Asp Phe Phe Ala Trp Ala Lys Val Asn Gln Val Pro Val
                                  250
Thr His Glu Phe Lys Val Pro Arg Glu Leu Ala Gly Thr Lys Gly Ala
                              265
Glu Lys Ser Leu Lys Arg Pro Leu Gly Asp Val Thr Asn Thr Ser Tyr
                          280
Lys Ser Leu Glu Lys Arg Ala Arg Leu Ser Phe Val Pro Glu Thr Pro
                     295
Arg Ser Ser Asp Val Thr Val Asp Pro Ala Pro Leu Arg Pro Leu Asn
                  310
                                      315
Trp Asn Ser Arg Tyr Asp Cys Lys Cys Asp Tyr His Ala Gln Phe Asp
                                  330
Asn Ile Ser Asn Lys Cys Asp Glu Cys Glu Tyr Leu Asn Arg Gly Lys
                              345
Asn Gly Cys Ile Cys His Asn Val Thr His Cys Gln Ile Cys His Gly
                          360
Ile Pro Pro Trp Glu Lys Glu Asn Leu Ser Asp Phe Gly Asp Phe Asp
                      375
Asp Ala Asn Lys Glu Gln
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     <211> 610
     <212> PRT
     <213> Artificial Sequence
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<223> Description of Artificial Sequence:/Note =

<220>

## synthetic construct

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His Gln Gln Pro Leu Glu Asp Arg Met Phe Lys Phe Glu Leu Thr Lys
                           440
Arg Leu Pro Pro Asp Phe Gly Lys Ile Thr Lys Gln Glu Val Lys Asp
                      455
                                          460
Phe Phe Ala Trp Ala Lys Val Asn Gln Val Pro Val Thr His Glu Phe
                  470
                                      475
Lys Val Pro Arg Glu Leu Ala Gly Thr Lys Gly Ala Glu Lys Ser Leu
               485
                                  490
Lys Arg Pro Leu Gly Asp Val Thr Asn Thr Ser Tyr Lys Ser Leu Glu
                              505
Lys Arg Ala Arg Leu Ser Phe Val Pro Glu Thr Pro Arg Ser Ser Asp
                      520
                                       525
Val Thr Val Asp Pro Ala Pro Leu Arg Pro Leu Asn Trp Asn Ser Arg
           535
                                         540
Tyr Asp Cys Lys Cys Asp Tyr His Ala Gln Phe Asp Asn Ile Ser Asn
                 550
                          555
Lys Cys Asp Glu Cys Glu Tyr Leu Asn Arg Gly Lys Asn Gly Cys Ile
              565
                                 570
Cys His Asn Val Thr His Cys Gln Ile Cys His Gly Ile Pro Pro Trp
                              585
Glu Lys Glu Asn Leu Ser Asp Phe Gly Asp Phe Asp Asp Ala Asn Lys
Glu Gln
   610
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     <211> 724
     <212> PRT
     <213> Artificial Sequence
     <223> Description of Artificial Sequence:/Note =
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Gly Leu Arg Glu Phe Leu Gly Leu Glu Ala Gly Pro Pro Lys Pro Lys
          20
                              25
Pro Asn Gln Gln His Gln Asp Gln Ala Arg Gly Leu Val Leu Pro Gly
                          40
Tyr Asn Tyr Leu Gly Pro Gly Asn Gly Leu Asp Arg Gly Glu Pro Val
                      -55
Asn Arg Ala Asp Glu Val Ala Arg Glu His Asp Ile Ser Tyr Asn Glu
Gln Leu Glu Ala Gly Asp Asn Pro Tyr Leu Lys Tyr Asn His Ala Asp
                                  90
Ala Glu Phe Gln Glu Lys Leu Ala Asp Asp Thr Ser Phe Gly Gly Asn
                              105
Leu Gly Lys Ala Val Phe Gln Ala Lys Lys Arg Val Leu Glu Pro Phe
                          120
Gly Leu Val Glu Glu Gly Ala Lys Thr Ala Pro Thr Gly Lys Arg Ile
                      135
                                          140
Asp Asp His Phe Pro Lys Arg Lys Lys Ala Arg Thr Glu Glu Asp Ser
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150

165

Lys Pro Ser Thr Ser Ser Asp Ala Glu Ala Gly Pro Ser Gly Ser Gln

155

Gln Leu Gln Ile Pro Ala Gln Pro Ala Ser Ser Leu Gly Ala Asp Thr Met Ser Ala Gly Gly Gly Pro Leu Gly Asp Asn Asn Gln Gly Ala Asp Gly Val Gly Asn Ala Ser Gly Asp Trp His Cys Asp Ser Thr Trp Met Gly Asp Arg Val Val Thr Lys Ser Thr Arg Thr Trp Val Leu Pro Ser Tyr Asn Asn His Gln Tyr Arg Glu Ile Lys Ser Gly Ser Val Asp Gly Ser Asn Ala Asn Ala Tyr Phe Gly Tyr Ser Thr Pro Trp Gly Tyr Phe Asp Phe Asn Arg Phe His Ser His Trp Ser Pro Arg Asp Trp Gln Arg Leu Ile Asn Asn Tyr Trp Gly Phe Arg Pro Arg Ser Leu Arg Val Lys Ile Phe Asn Ile Gln Val Lys Glu Val Thr Val Gln Asp Ser Thr Thr Thr Ile Ala Asn Asn Leu Thr Ser Thr Val Gln Val Phe Thr Asp Asp Asp Tyr Gln Leu Pro Tyr Val Val Gly Asn Gly Thr Glu Gly Cys Leu Pro Ala Phe Pro Pro Gln Val Phe Thr Leu Pro Gln Tyr Gly Tyr Ala Thr Leu Asn Arg Asp Asn Thr Glu Asn Pro Thr Glu Arg Ser Ser Phe Phe Cys Leu Glu Tyr Phe Pro Ser Lys Met Leu Arg Thr Gly Asn Asn Phe Glu Phe Thr Tyr Asn Phe Glu Glu Val Pro Phe His Ser Ser Phe Ala Pro Ser Gln Asn Leu Phe Lys Leu Ala Asn Pro Leu Val Asp Gln Tyr Leu Tyr Arg Phe Val Ser Thr Asn Asn Thr Gly Gly Val Gln Phe Asn Lys Asn Leu Ala Gly Arg Tyr Ala Asn Thr Tyr Lys Asn Trp Phe Pro Gly Pro Met Gly Arg Thr Gln Gly Trp Asn Leu Gly Ser Gly Val Asn Arg Ala Ser Val Ser Ala Phe Ala Thr Thr Asn Arg Met Glu Leu Glu Gly Ala Ser Tyr Gln Val Pro Pro Gln Pro Asn Gly Met Thr Asn Asn Leu Gln Gly Ser Asn Thr Tyr Ala Leu Glu Asn Thr Met Ile Phe Asn Ser Gln Pro Ala Asn Pro Gly Thr Thr Ala Thr Tyr Leu Glu Gly Asn Met Leu Ile Thr Ser Glu Ser Glu Thr Gln Pro Val Asn Arg Val Ala Tyr Asn Val Gly Gly Gln Met Ala Thr Asn Asn Gln Ser Ser Thr Thr Ala Pro Ala Thr Gly Thr Tyr Asn Leu Gln Glu Ile Val Pro Gly Ser Val Trp Met Glu Arg Asp Val Tyr Leu Gln Gly Pro Ile Trp Ala Lys Ile Pro Glu Thr Gly Ala His Phe His Pro Ser Pro Ala Met Gly Gly Phe Gly Leu Lys His Pro Pro Pro Met Met Leu Ile Lys Asn 

7

Thr Pro Val Pro Gly Asn Ile Thr Ser Phe Ser Asp Val Pro Val Ser 655

Ser Phe Ile Thr Gln Tyr Ser Thr Gly Gln Val Thr Val Glu Met Glu 660

Trp Glu Leu Lys Lys Glu Asn Ser Lys Arg Trp Asn Pro Glu Ile Gln 670

Tyr Thr Asn Asn Tyr Asn Asp Pro Gln Phe Val Asp Phe Ala Pro Asp 690

Ser Thr Gly Glu Tyr Arg Thr Arg Pro Ile Gly Thr Arg Tyr Leu 720

Thr Arg Pro Leu

<210> 5

<211> 588

<212> PRT

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:/Note = synthetic construct

<400> 5

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Glu Glu Val Pro Phe His Ser Ser Phe Ala Pro Ser Gln Asn Leu Phe
                          280
Lys Leu Ala Asn Pro Leu Val Asp Gln Tyr Leu Tyr Arg Phe Val Ser
                      295
                                         300
Thr Asn Asn Thr Gly Gly Val Gln Phe Asn Lys Asn Leu Ala Gly Arg
                  310
                           315
Tyr Ala Asn Thr Tyr Lys Asn Trp Phe Pro Gly Pro Met Gly Arg Thr
              325
                           330
Gln Gly Trp Asn Leu Gly Ser Gly Val Asn Arg Ala Ser Val Ser Ala
                             345
Phe Ala Thr Thr Asn Arg Met Glu Leu Glu Gly Ala Ser Tyr Gln Val
                   360
                                            365
Pro Pro Gln Pro Asn Gly Met Thr Asn Asn Leu Gln Gly Ser Asn Thr
                375
                              380
Tyr Ala Leu Glu Asn Thr Met Ile Phe Asn Ser Gln Pro Ala Asn Pro
            390
                           395
Gly Thr Thr Ala Thr Tyr Leu Glu Gly Asn Met Leu Ile Thr Ser Glu
              405
                                410
Ser Glu Thr Gln Pro Val Asn Arg Val Ala Tyr Asn Val Gly Gln
          420
                             425
Met Ala Thr Asn Asn Gln Ser Ser Thr Thr Ala Pro Ala Thr Gly Thr
                          440
Tyr Asn Leu Gln Glu Ile Val Pro Gly Ser Val Trp Met Glu Arg Asp
                      455
Val Tyr Leu Gln Gly Pro Ile Trp Ala Lys Ile Pro Glu Thr Gly Ala
                  470
                                     475
His Phe His Pro Ser Pro Ala Met Gly Gly Phe Gly Leu Lys His Pro
               485
                                 490
Pro Pro Met Met Leu Ile Lys Asn Thr Pro Val Pro Gly Asn Ile Thr
           500
                             505
Ser Phe Ser Asp Val Pro Val Ser Ser Phe Ile Thr Gln Tyr Ser Thr
                         520
                                             525
Gly Gln Val Thr Val Glu Met Glu Trp Glu Leu Lys Lys Glu Asn Ser
                      535
                                        540
Lys Arg Trp Asn Pro Glu Ile Gln Tyr Thr Asn Asn Tyr Asn Asp Pro
                  550
                                     555
Gln Phe Val Asp Phe Ala Pro Asp Ser Thr Gly Glu Tyr Arg Thr Thr
              565
                           570
Arg Pro Ile Gly Thr Arg Tyr Leu Thr Arg Pro Leu
           580
     <210> 6
     <211> 532
     <212> PRT
     <213> Artificial Sequence
     <223> Description of Artificial Sequence:/Note =
           synthetic construct
     <400> 6
Met Ser Ala Gly Gly Gly Pro Leu Gly Asp Asn Asn Gln Gly Ala
                                 10
Asp Gly Val Gly Asn Ala Ser Gly Asp Trp His Cys Asp Ser Thr Trp
                             25
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Met Gly Asp Arg Val Val Thr Lys Ser Thr Arg Thr Trp Val Leu Pro 35 40 45

Ser Tyr Asn Asn His Gln Tyr Arg Glu Ile Lys Ser Gly Ser Val Asp Gly Ser Asn Ala Asn Ala Tyr Phe Gly Tyr Ser Thr Pro Trp Gly Tyr 70 75 Phe Asp Phe Asn Arg Phe His Ser His Trp Ser Pro Arg Asp Trp Gln 8.5 90 Arg Leu Ile Asn Asn Tyr Trp Gly Phe Arg Pro Arg Ser Leu Arg Val 105 Lys Ile Phe Asn Ile Gln Val Lys Glu Val Thr Val Gln Asp Ser Thr 120 Thr Thr Ile Ala Asn Asn Leu Thr Ser Thr Val Gln Val Phe Thr Asp 135 140 Asp Asp Tyr Gln Leu Pro Tyr Val Val Gly Asn Gly Thr Glu Gly Cys 150 155 Leu Pro Ala Phe Pro Pro Gln Val Phe Thr Leu Pro Gln Tyr Gly Tyr 165 170 Ala Thr Leu Asn Arg Asp Asn Thr Glu Asn Pro Thr Glu Arg Ser Ser 185 Phe Phe Cys Leu Glu Tyr Phe Pro Ser Lys Met Leu Arg Thr Gly Asn 200 Asn Phe Glu Phe Thr Tyr Asn Phe Glu Glu Val Pro Phe His Ser Ser 215 Phe Ala Pro Ser Gln Asn Leu Phe Lys Leu Ala Asn Pro Leu Val Asp 230 235 Gln Tyr Leu Tyr Arg Phe Val Ser Thr Asn Asn Thr Gly Gly Val Gln 245 250 Phe Asn Lys Asn Leu Ala Gly Arg Tyr Ala Asn Thr Tyr Lys Asn Trp 265 Phe Pro Gly Pro Met Gly Arg Thr Gln Gly Trp Asn Leu Gly Ser Gly 280 Val Asn Arg Ala Ser Val Ser Ala Phe Ala Thr Thr Asn Arg Met Glu 295 Leu Glu Gly Ala Ser Tyr Gln Val Pro Pro Gln Pro Asn Gly Met Thr 310 315 Asn Asn Leu Gln Gly Ser Asn Thr Tyr Ala Leu Glu Asn Thr Met Ile 325 330 Phe Asn Ser Gln Pro Ala Asn Pro Gly Thr Thr Ala Thr Tyr Leu Glu 345 Gly Asn Met Leu Ile Thr Ser Glu Ser Glu Thr Gln Pro Val Asn Arg 360 Val Ala Tyr Asn Val Gly Gly Gln Met Ala Thr Asn Asn Gln Ser Ser 375 380 Thr Thr Ala Pro Ala Thr Gly Thr Tyr Asn Leu Gln Glu Ile Val Pro 390 395 Gly Ser Val Trp Met Glu Arg Asp Val Tyr Leu Gln Gly Pro Ile Trp 405 410 Ala Lys Ile Pro Glu Thr Gly Ala His Phe His Pro Ser Pro Ala Met 420 425 Gly Gly Phe Gly Leu Lys His Pro Pro Pro Met Met Leu Ile Lys Asn 440 Thr Pro Val Pro Gly Asn Ile Thr Ser Phe Ser Asp Val Pro Val Ser 455 460 Ser Phe Ile Thr Gln Tyr Ser Thr Gly Gln Val Thr Val Glu Met Glu 470 475 Trp Glu Leu Lys Lys Glu Asn Ser Lys Arg Trp Asn Pro Glu Ile Gln 485 490 Tyr Thr Asn Asn Tyr Asn Asp Pro Gln Phe Val Asp Phe Ala Pro Asp 500 505

10

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Ser Thr Gly Glu Tyr Arg Thr Thr Arg Pro Ile Gly Thr Arg Tyr Leu
        515
                            520
Thr Arg Pro Leu
    530
      <210> 7
      <211> 2307
      <212> DNA
      <213> Artificial Sequence
      <220>
      <223> Description of Artificial Sequence:/Note =
            synthetic construct
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                                                                        60
ctgcgaccgc tcaattggaa ttcaagtaaa taaagcgagt agtcatgtct tttgttgatc
                                                                       120
accetecaga ttggttggaa gaagttggtg aaggtetteg egagtttttg ggeettgaag
                                                                       180
cgggcccacc gaaaccaaaa cccaatcagc agcatcaaga tcaagcccgt ggtcttgtgc
                                                                       240
tgcctggtta taactatctc ggacccggaa acggtctcga tcgaggagag cctgtcaaca
                                                                       300
gggcagacga ggtcgcgcga gagcacgaca tctcgtacaa cgagcagctt gaggcgggag
                                                                       360
acaaccccta cctcaagtac aaccacgcgg acgccgagtt tcaggagaag ctcgccgacg
                                                                       420
acacatcctt cgggggaaac ctcggaaagg cagtctttca ggccaagaaa agggttctcg
                                                                       480
aaccttttgg cctggttgaa gagggtgcta agacggcccc taccggaaag cggatagacg
                                                                       540
accactttcc aaaaagaaag aaggctcgga ccgaagagga ctccaagcct tccacctcgt
                                                                       600
cagacgccga agctggaccc agcggatccc agcagctgca aatcccagcc caaccagcct
                                                                       660
caagtttggg agctgataca atgtctgcgg gaggtggcgg cccattgggc gacaataacc
                                                                       720
aaggtgccga tggagtgggc aatgcctcgg gagattggca ttgcgattcc acgtggatqg
                                                                       780
gggacagagt cgtcaccaag tccacccgaa cctgggtgct gcccagctac aacaaccacc
                                                                       840
agtaccgaga gatcaaaagc ggctccgtcg acggaagcaa cgccaacgcc tactttggat
                                                                       900
acagcacccc ctgggggtac tttgacttta accgcttcca cagccactgg agcccccgag
                                                                       960
actggcaaag actcatcaac aactactggg gcttcagacc ccggtccctc agagtcaaaa
                                                                      1020
tetteaacat teaagteaaa gaggteaegg tgeaggaete caccaccace ategeeaaca
                                                                      1080
acctcacctc caccgtccaa gtgtttacgg acgacgacta ccagctgccc tacgtcgtcg
                                                                      1140
gcaacgggac cgagggatgc ctgccggcct tccctccgca ggtctttacg ctgccgcagt
                                                                      1200
acggttacgc gacgctgaac cgcgacaaca cagaaaatcc caccgagagg agcagcttct
                                                                     1260
tctgcctaga gtactttccc agcaagatgc tgagaacggg caacaacttt gagtttacct
                                                                     1320
acaactttga ggaggtgccc ttccactcca gcttcgctcc cagtcagaac ctgttcaagc
                                                                     1380
tggccaaccc gctggtggac cagtacttgt accgcttcgt gagcacaaat aacactqqcq
                                                                     1440
gagtccagtt caacaagaac ctggccggga gatacgccaa cacctacaaa aactqqttcc
                                                                     1500
cggggcccat gggccgaacc cagggctgga acctgggctc cggggtcaac cgcqccaqtq
                                                                     1560
tcagcgcctt cgccacgacc aataggatgg agctcgaggg cgcgagttac caggtgcccc
                                                                     1620
cgcagccgaa cggcatgacc aacaacctcc agggcagcaa cacctatgcc ctggagaaca
                                                                     1680
ctatgatett caacagecag ceggegaace egggeaceae egecaegtae eteqaqqqea
                                                                     1740
acatgeteat caccagegag agegagaege ageeggtgaa eegegtggeg tacaaegteg
                                                                     1800
gcgggcagat ggccaccaac aaccagagct ccaccactgc ccccgcgacc ggcacgtaca
                                                                     1860
acctccagga aatcgtgccc ggcagcgtgt ggatggagag ggacgtgtac ctccaaggac
                                                                     1920
ccatctgggc caagatccca gagacggggg cgcactttca cccctctccg gccatgggcg
                                                                     1980
gatteggaet caaacaccca eegeceatga tgeteateaa gaacaegeet gtgeeeggaa
                                                                     2040
atatcaccag cttctcggac gtgcccgtca gcagcttcat cacccagtac agcaccgggc
                                                                     2100
aggtcaccgt ggagatggag tgggagctca agaaggaaaa ctccaagagg tggaacccag
                                                                     2160
agatccagta cacaaacaac tacaacgacc cccagtttgt ggactttgcc ccggacagca
                                                                     2220
ccggggaata cagaaccacc agacctatcg gaacccgata ccttacccga cccctttaac
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ccattcatgt cgcataccct caataaa
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      <210> 8
      <211> 2264
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<212> DNA

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<213> Artificial Sequence
      <220>
      <223> Description of Artificial Sequence:/Note =
            synthetic construct
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                                                                       120
gtttttgggc cttgaagcgg gcccaccgaa accaaaaccc aatcagcagc atcaagatca
                                                                       180
agcccgtggt cttgtgctgc ctggttataa ctatctcgga cccggaaacg gtctcgatcg
                                                                       240
aggagagect gtcaacaggg cagacgaggt cgcgcgagag cacgacatct cgtacaacga
                                                                       300
gcagcttgag gcgggagaca acccctacct caagtacaac cacgcggacg ccgagtttca
                                                                       360
ggagaagete geegaegaea cateettegg gggaaacete ggaaaggeag tettteagge
                                                                       420
caagaaaagg gttctcgaac cttttggcct ggttgaagag ggtgctaaqa cqqccctac
                                                                       480
cggaaagcgg atagacgacc actttccaaa aagaaagaag gctcggaccg aagaggactc
                                                                       540
caagcettee acctegteag acgeegaage tggacecage ggateecage agetgeaaat
                                                                       600
cccagcccaa ccagcctcaa gtttgggagc tgatacaatg tctgcgggag gtggcgqccc
                                                                       660
attgggcgac aataaccaag gtgccgatgg agtgggcaat gcctcgggag attggcattg
                                                                       720
cgattccacg tggatggggg acagagtcgt caccaagtcc acccgaacct gggtgctgcc
                                                                       780
cagctacaac aaccaccagt accgagagat caaaagcggc tccgtcgacg gaagcaacgc
                                                                       840
caacgcctac tttggataca gcacccctg ggggtacttt gactttaacc gcttccacag
                                                                       900
ccactggagc ccccgagact ggcaaagact catcaacaac tactggggct tcagaccccg
                                                                       960
gtccctcaga gtcaaaatct tcaacattca agtcaaagag gtcacggtgc aggactccac
                                                                      1020
caccaccatc gccaacaacc tcacctccac cgtccaagtg tttacggacg acgactacca
                                                                      1080
gctgccctac gtcgtcggca acgggaccga gggatgcctg ccggccttcc ctccgcaggt
                                                                     1140
ctttacgctg ccgcagtacg gttacgcgac gctgaaccgc gacaacacag aaaatcccac
                                                                      1200
cgagaggagc agcttcttct gcctagagta ctttcccagc aagatgctga gaacgggcaa
                                                                      1260
caactttgag tttacctaca actttgagga ggtgcccttc cactccagct tcgctcccag
                                                                      1320
tragaarctg ttraagetgg craacceget ggtggarcag tarttgtare gettegtgag
                                                                      1380
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cacgtacctc gagggcaaca tgctcatcac cagcgagagc gagacgcagc cggtgaaccg
                                                                     1740
cgtggcgtac aacgtcggcg ggcagatggc caccaacaac cagagctcca ccactgcccc
                                                                     1800
cgcgaccggc acgtacaacc tccaggaaat cgtgcccggc agcgtgtgga tggagaggga
                                                                     1860
cgtgtacctc caaggaccca tctgggccaa gatcccagag acgggggcgc actttcaccc
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ctctccggcc atgggcggat tcggactcaa acacccaccg cccatgatgc tcatcaagaa
                                                                     1980
cacgcetgtg cccggaaata tcaccagctt ctcggacgtg cccgtcagca gcttcatcac
                                                                     2040
ccagtacagc accgggcagg tcaccgtgga gatggagtgg gagctcaaga aggaaaactc
                                                                     2100
caagaggtgg aacccagaga tccagtacac aaacaactac aacgaccccc agtttgtgga
                                                                     2160
ctttgccccg gacagcaccg gggaatacag aaccaccaga cctatcggaa cccgatacct
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                                                                     2264
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            synthetic construct
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gtttttgggc cttgaagcgg gcccaccgaa a	accaaaaccc	aatcagcagc	atcaagatca	180
agcccgtggt cttgtgctgc ctggttataa o	ctatctcgga	cccggaaacg	gtctcgatcg	240
aggagageet gteaacaggg cagaegaggt o	cgcgcgagag	cacgacatct	cgtacaacga	300
gcagettgag gegggagaea acceetaeet o	caagtacaac	cacgcggacg	ccgagtttca	360
ggagaagete geegaegaea eateettegg g	gggaaacctc	ggaaaggcag	tctttcaggc	420
caagaaaagg gttctcgaac cttttggcct g	ggttgaagag	ggtgctaaga	cggcccctac	480
cggaaagcgg atagacgacc actttccaaa a	aagaaagaag	gctcggaccg	aagaggactc	540
caagcettee acetegteag aegeegaage t	tggacccagc	ggatcccagc	agctgcaaat	600
cccagcccaa ccagcctcaa gtttgggagc t	tgatacaatg	tctgcgggag	gtggcggccc	660
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gtccctcaga gtcaaaatct tcaacattca a	agtcaaagag	gtcacggtgc	aggactccac	1020
caccaccatc gccaacaacc tcacctccac o	cgtccaagtg	tttacggacg	acgactacca	1080
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ggtcaaccgc gccagtgtca gcgccttcgc c	cacgaccaat	aggatggagc	tcgagggcgc	1560
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cgcgaccggc acgtacaacc tccaggaaat c				1860
cgtgtacctc caaggaccca tctgggccaa g				1920
ctctccggcc atgggcggat tcggactcaa a				1980
cacgectgtg ceeggaaata teaccagett c	ratogacycy	gaggtgagga	getteateae	2040
ccagtacagc accgggcagg tcaccgtgga g caagaggtgg aacccagaga tccagtacac a	acggagtgg	ageceaaga	aggaaaactc	2100
ctttgccccg gacagcaccg gggaatacag a	aaacaaccac	cctatcccaa	ageregega	2160
taccegacce ctttaaccca ttcatgtege a			eeegataeet	2220
accegace eccaacea cecaegeege a	acaccccaa	caaa		2264
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<213> Artificial Sequence				
<220>				
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synthetic construct	1	,		
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tggcgctcgt caactggctc gtggagcacg g	gcatcacttc	cgagaagcag	tggatccaga	180
aaaatcagga gagctacctc tccttcaact c	ccaccggcaa	ctctcggaqc	cagatcaagg	240
ccgcgctcga caacgcgacc aaaattatga g	gtctgacaaa	aagcgcggtq	gactacctca	300
tggggagctc cgttcccgag gacatttcaa a	aaacagaat	ctggcaaatt	tttgagatga	360
atggctacga cccggcctac gcgggatcca t	tcctctacgg	ctggtgtcag	cgctccttca	420
acaagaggaa caccgtctgg ctctacggac c	ccgccacgac	cggcaagacc	aacatcgcgg	480
aggccatcgc ccacactgtg cccttttacg g	gctgcgtgaa	ctggaccaat	gaaaactttc	540
cctttaatga ctgtgtggac aaaatgctca t	tttggtggga	ggagggaaag	atgaccaaca	600
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13

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gtgtggtggt ggatgggaat tccacgacct ttgaacacca gcagccgctg gaggaccgca
                                                                       780
tgttcaaatt tgaactgact aagcggctcc cgccagattt tggcaagatt actaagcagg
                                                                       840
aagtcaagga cttttttgct tgggcaaagg tcaatcaggt gccggtgact cacgagttta
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aagttcccag ggaattggcg ggaactaaag gggcggagaa atctctaaaa cgcccactgg
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gtgacgtcac caatactagc tataaaagtc tggagaagcg ggccaggctc tcatttgttc
                                                                      1020
ccgagacgcc tcgcagttca gacgtgactg ttgatcccgc tcctctgcga ccgctcaatt
                                                                      1080
ggaattcaag gtatgattgc aaatgtgact atcatgctca atttgacaac atttctaaca
                                                                      1140
aatgtgatga atgtgaatat ttgaatcggg gcaaaaatgg atgtatctgt cacaatgtaa
                                                                      1200
ctcactgtca aatttgtcat gggattcccc cctgggaaaa ggaaaacttg tcagattttg
                                                                      1260
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                                                                      1292
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      <213> Artificial Sequence
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            synthetic construct
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                                                                       120
ctgggtaact ggtcaaattt gggagctgcc tccagagtca gatttaaatt tqactctgqt
                                                                       180
tgaacagcct cagttgacgg tggctgatag aattcgccgc gtgttcctgt acgagtggaa
                                                                       240
caaattttcc aagcaggagt ccaaattctt tgtgcagttt gaaaagggat ctgaatattt
                                                                       300
tcatctgcac acgcttgtgg agacctccgg catctcttcc atggtcctcg gccgctacgt
                                                                       360
gagtcagatt cgcgcccagc tggtgaaagt ggtcttccag ggaattgaac cccagatcaa
                                                                       420
cgactgggtc gccatcacca aggtaaagaa gggcggagcc aataaggtgg tggattctgg
                                                                       480
gtatattccc gcctacctgc tgccgaaggt ccaaccggag cttcagtggg cgtggacaaa
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                                                                       660
tgacccggtc atcaaaagca agacttccca gaaatacatg gcgctcgtca actggctcgt
                                                                      720
ggagcacggc atcacttccg agaagcagtg gatccaggaa aatcaggaga gctacctctc
                                                                      780
cttcaactcc accggcaact ctcggagcca gatcaaggcc gcgctcgaca acqcqaccaa
                                                                       840
aattatgagt ctgacaaaaa gcgcggtgga ctacctcgtg gggagctccg ttcccgagga
                                                                      900
catttcaaaa aacagaatct ggcaaatttt tgagatgaat ggctacgacc cggcctacqc
                                                                      960
gggatecate etetaegget ggtgteageg eteetteaac aagaggaaca eegtetgget
                                                                     1020
ctacggaccc gccacgaccg gcaagaccaa catcgcggag gccatcgccc acactgtgcc
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cttttacggc tgcgtgaact ggaccaatga aaactttccc tttaatgact gtgtggacaa
                                                                     1140
aatgctcatt tggtgggagg agggaaagat gaccaacaag gtggttgaat ccgccaaggc
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catcctgggg ggctcaaagg tgcgggtcga tcagaaatgt aaatcctctg ttcaaattga
                                                                     1260
ttctacccct gtcattgtaa cttccaatac aaacatgtgt gtggtggtgg atgggaattc
                                                                     1320
cacgacettt gaacaccage ageegetgga ggacegeatg ttcaaatttg aactgactaa
                                                                     1380
geggeteeeg ceagattttg geaagattae taageaggaa gteaaggaet tttttgettg
                                                                     1440
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                                                                     1500
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                                                                     1560
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                                                                     1620
cgtgactgtt gatcccgctc ctctgcgacc gctcaattgg aattcaaggt atgattgcaa
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                                                                     1740
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gattccccc tgggaaaagg aaaacttgtc agattttggg gattttgacg atgccaataa
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agaacagtaa
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<211> 330

<212> PRT

14

<213> Artificial Sequence

<220>

<223> Description of Artificial Sequence:/Note = synthetic construct

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<210> 13

<211> 1115

<212> DNA

<213> Artificial Sequence

Trp Asn Ser Arg Leu Val Gly Arg Ser Trp 325

<223> Description of Artificial Sequence:/Note = synthetic construct

15

195

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                                                                       120
acatggeget egteaactgg etegtggage aeggeateae tteegagaag eaqtqqatee
                                                                       180
aggaaaatca ggagagctac ctctccttca actccaccgg caactctcgg agccagatca
                                                                       240
aggccgcgct cgacaacgcg accaaaatta tgagtctgac aaaaagcgcg gtggactacc
                                                                       300
tcgtggggag ctccgttccc gaggacattt caaaaaacag aatctggcaa atttttgaga
                                                                       360
tgaatggcta cgacccggcc tacgcgggat ccatcctcta cggctggtgt cagcgctcct
                                                                       420
tcaacaagag gaacaccgtc tggctctacg gacccgccac gaccggcaag accaacatcg
                                                                       480
eggaggeeat egeceacaet gtgeeetttt aeggetgegt gaaetggaee aatgaaaaet
                                                                       540
ttccctttaa tgactgtgtg gacaaaatgc tcatttggtg ggaggaggga aagatgacca
                                                                       600
acaaggtggt tgaatccgcc aaggccatcc tggggggctc aaaggtgcgg gtcgatcaga
                                                                       660
aatgtaaatc ctctgttcaa attgattcta cccctgtcat tgtaacttcc aatacaaaca
                                                                       720
tgtgtgtggt ggtggatggg aattccacga cctttgaaca ccagcagccg ctqqaqqacc
                                                                       780
gcatgttcaa atttgaactg actaagcggc tcccgccaga ttttqqcaaq attactaaqc
                                                                       840
aggaagtcaa ggactttttt gcttgggcaa aggtcaatca ggtgccggtg actcacqaqt
                                                                       900
ttaaagttcc cagggaattg gcgggaacta aaggggcgga gaaatctcta aaacqcccac
                                                                       960
tqqqtgacgt caccaatact agctataaaa gtctggagaa gcgggccagg ctctcatttg
                                                                      1020
ttcccgagac gcctcgcagt tcagacgtga ctgttgatcc cgctcctctg cgaccgctca
                                                                      1080
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                                                                      1115
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      <211> 550
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      <223> Description of Artificial Sequence:/Note =
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Glu His Leu Pro Gly Ile Ser Asp Ser Phe Val Asp Trp Val Thr Gly
                                25
Gln Ile Trp Glu Leu Pro Pro Glu Ser Asp Leu Asn Leu Thr Leu Val
                            40
Glu Gln Pro Gln Leu Thr Val Ala Asp Arg Ile Arg Arg Val Phe Leu
Tyr Glu Trp Asn Lys Phe Ser Lys Gln Glu Ser Lys Phe Phe Val Gln
                    70
                                        75
Phe Glu Lys Gly Ser Glu Tyr Phe His Leu His Thr Leu Val Glu Thr
                8.5
                                    90
Ser Gly Ile Ser Ser Met Val Leu Gly Arg Tyr Val Ser Gln Ile Arg
                                105
Ala Gln Leu Val Lys Val Val Phe Gln Gly Ile Glu Pro Gln Ile Asn
                            120
Asp Trp Val Ala Ile Thr Lys Val Lys Lys Gly Gly Ala Asn Lys Val
                        135
                                            140
Val Asp Ser Gly Tyr Ile Pro Ala Tyr Leu Leu Pro Lys Val Gln Pro
                    150
                                        155
Glu Leu Gln Trp Ala Trp Thr Asn Leu Asp Glu Tyr Lys Leu Ala Ala
                165
                                    170
Leu Asn Leu Glu Glu Arg Lys Arg Leu Val Ala Gln Phe Leu Ala Glu
                                185
Ser Ser Gln Arg Ser Gln Glu Ala Ala Ser Gln Arg Glu Phe Ser Ala
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200

16

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Asp Pro Val Ile Lys Ser Lys Thr Ser Gln Lys Tyr Met Ala Leu Val
                        215
                                            220
Asn Trp Leu Val Glu His Gly Ile Thr Ser Glu Lys Gln Trp Ile Gln
                    230
                                        235
Glu Asn Gln Glu Ser Tyr Leu Ser Phe Asn Ser Thr Gly Asn Ser Arg
                245
                                    250
Ser Gln Ile Lys Ala Ala Leu Asp Asn Ala Thr Lys Ile Met Ser Leu
            260
                                265
Thr Lys Ser Ala Val Asp Tyr Leu Val Gly Ser Ser Val Pro Glu Asp
                           280
                                                285
Ile Ser Lys Asn Arg Ile Trp Gln Ile Phe Glu Met Asn Gly Tyr Asp
                       295
                                           300
Pro Ala Tyr Ala Gly Ser Ile Leu Tyr Gly Trp Cys Gln Arg Ser Phe
                   310
                                       315
Asn Lys Arg Asn Thr Val Trp Leu Tyr Gly Pro Ala Thr Thr Gly Lys
               325
                                   330
Thr Asn Ile Ala Glu Ala Ile Ala His Thr Val Pro Phe Tyr Gly Cys
            340
                               345
Val Asn Trp Thr Asn Glu Asn Phe Pro Phe Asn Asp Cys Val Asp Lys
                           360
Met Leu Ile Trp Trp Glu Glu Gly Lys Met Thr Asn Lys Val Val Glu
                        375
Ser Ala Lys Ala Ile Leu Gly Gly Ser Lys Val Arg Val Asp Gln Lys
                    390
                                        395
Cys Lys Ser Ser Val Gln Ile Asp Ser Thr Pro Val Ile Val Thr Ser
                405
                                    410
Asn Thr Asn Met Cys Val Val Val Asp Gly Asn Ser Thr Thr Phe Glu
                                425
His Gln Gln Pro Leu Glu Asp Arg Met Phe Lys Phe Glu Leu Thr Lys
                            440
Arg Leu Pro Pro Asp Phe Gly Lys Ile Thr Lys Gln Glu Val Lys Asp
                        455
                                            460
Phe Phe Ala Trp Ala Lys Val Asn Gln Val Pro Val Thr His Glu Phe
                    470
                                        475
Lys Val Pro Arg Glu Leu Ala Gly Thr Lys Gly Ala Glu Lys Ser Leu
                485
                                   490
Lys Arg Pro Leu Gly Asp Val Thr Asn Thr Ser Tyr Lys Ser Leu Glu
            500
                               505
Lys Arg Ala Arg Leu Ser Phe Val Pro Glu Thr Pro Arg Ser Ser Asp
                           520
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Val Thr Val Asp Pro Ala Pro Leu Arg Pro Leu Asn Trp Asn Ser Arg
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Leu Val Gly Arg Ser Trp
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      <223> Description of Artificial Sequence:/Note =
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ctgggtaact ggtcaaattt gggagctgcc tccagagtca gatttaaatt tgactctggt

17

tgaacagcct cagttgacgg tggctgatag aattcgccgc gtgttcctgt acgagtggaa 240 caaattttcc aagcaggagt ccaaattctt tgtgcagttt gaaaagggat ctqaatattt 300 tcatctgcac acgcttgtgg agacctccgg catctcttcc atggtcctcg gccgctacgt 360 gagtcagatt cgcgcccagc tggtgaaagt ggtcttccag ggaattgaac cccagatcaa 420 cgactgggtc gccatcacca aggtaaagaa gggcggagcc aataaggtgg tggattctgg 480 gtatattccc gcctacctgc tgccgaaggt ccaaccggag cttcagtggg cgtggacaaa 540 cctggacgag tataaattgg ccgccctgaa tctggaggag cgcaaacggc tcgtcgcgca 600 gtttctggca gaatcctcgc agcgctcgca ggaggcggct tcgcagcgtg agttctcggc 660 tgacccggtc atcaaaagca agacttccca gaaatacatg gcgctcgtca actggctcgt 720 ggagcacggc atcacttccg agaagcagtg gatccaggaa aatcaggaga gctacctctc 780 cttcaactcc accggcaact ctcggagcca gatcaaggcc gcgctcgaca acgcgaccaa 840 aattatgagt ctgacaaaaa gcgcggtgga ctacctcgtg gggagctccg ttcccqaqqa 900 catttcaaaa aacagaatct ggcaaatttt tgagatgaat ggctacgacc cqqcctacqc 960 gggatccatc ctctacggct ggtgtcagcg ctccttcaac aagaggaaca ccgtctqqct 1020 ctacggaccc gccacgaccg gcaagaccaa catcgcggag gccatcgccc acactgtgcc 1080 cttttacggc tgcgtgaact ggaccaatga aaactttccc tttaatgact gtgtggacaa 1140 aatgctcatt tggtgggagg agggaaagat gaccaacaag gtggttgaat ccgccaaggc 1200 catcctgggg ggctcaaagg tgcgggtcga tcagaaatgt aaatcctctg ttcaaattga 1260 ttctacccct gtcattgtaa cttccaatac aaacatgtgt gtggtggtgg atgggaattc 1320 cacgaccttt gaacaccagc agccgctgga ggaccgcatg ttcaaatttg aactgactaa 1380 gcggctcccg ccagattttg gcaagattac taagcaggaa gtcaaggact tttttgcttg 1440 ggcaaaggtc aatcaggtgc cggtgactca cgagtttaaa gttcccaggg aattggcggg 1500 aactaaaggg gcggagaaat ctctaaaacg cccactgggt gacgtcacca atactagcta 1560 taaaagtctg gagaagcggg ccaggctctc atttgttccc gagacgcctc gcagttcaga 1620 cgtgactgtt gatcccgctc ctctgcgacc gctcaattgg aattcaagat tggttggaag 1680 aagttggtga 1690 <210> 16 <211> 145 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:/Note = synthetic construct <400> 16 ccatcaccaa ggtaaagaag ggcggagcca ataaggtggt ggattctggg tatattcccg 60 cctacctgct gccgaaggtc caaccggagc ttcagtgggc gtggacaaac ctggacgagt 120 ataaattggc cgccctgaat ctgga 145 <210> 17 <211> 174 <212> DNA <213> Artificial Sequence <220> <223> Description of Artificial Sequence:/Note = synthetic construct <400> 17 taagcaggaa gtcaaggact tttttgcttg ggcaaaggtc aatcaggtgc cqqtqactca 60 cgagtttaaa gttcccaggg aattggcggg aactaaaggg gcggagaaat ctctaaaacq 120 cccactgggt gacgtcacca atactagcta taaaagtctg gagaaqcqqq ccaq 174 <210> 18

<211> 187 <212> DNA

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acgcgatagt taatgattaa cagtcatgtg atgtgtttta tccaatagga agaaagcgcg
                                                                       120
cgtatgagtt ctcgcgagac ttccggggta taaaagaccg agtgaacgag cccgccgcca
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ttctttq
                                                                       187
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tcgtttgggg gggtggcagc tcaaagagct gccagacgac ggccctctgg ccgtcgccc
                                                                       120
cccaaacgag ccagcgagcg agcgaacgcg acagggggga gagtgcca
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      <211> 168
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                                                                        60
tegtttgggg gggegaegge cagagggeeg tegtetgeeg getetttgag etgeeaecee
                                                                       120
cccaaacgag ccagcgagcg agcgaacgcg acagggggga gagtgcca
                                                                       168
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      <400> 21
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                                                                         8
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<223> Description of Artificial Sequence:/Note = synthetic construct

<400> 22

cggttgag

8

<210> 23

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<220>

<400> 23

caaaacctcc ttgcttgaga g